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(54) Title: DETECTION OF A NUCLEIC ACID SEQUENCE OR A CHANGE THEREIN

(57) Abstract

A method for detecting a specific polynucleotide sequence in a material is disclosed. The method includes exposing said material to an oligonucleotide primer having a sequence complementary to part of said specific polynucleotide sequence wherein said primer binds to part of said polynucleotide sequence when present in said material. Primer bound to the polynucleotide sequence is extended wherein any extended primer includes a detectable element and/or a separation element. Any extended primer is then separated into a fraction wherein said fraction does not have detectable element not included in said extended primer. One then determines whether any extended primer is present in said fraction by assaying said fraction for said extended primer wherein the presence of said extended primer is indicative of the presence of the specific polynucleotide sequence in said material and the absence of said extended primer in said fraction is indicative of the absence of the specific polynucleotide sequence in said material.

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Detection of a Nucleic Acid Sequence or a Change Therein.

Technical Field

This invention relates to a method for detecting a specific polynucleotide sequence in a material, a method for detecting at least two different specific polynucleotide sequences in material having a plurality of different polynucleotide sequences, methods for detecting whether a specific nucleotide or base is at a particular position in a specific polynucleotide sequence, methods for detecting whether the same or different specific nucleotides or bases are at particular positions in at least two different specific polynucleotide sequences in material having a plurality of different polynucleotide sequences, a screening method for detecting the presence of at least two different specific polynucleotide sequences in material having a plurality of different polynucleotide sequences and screening methods for detecting whether the same or different specific nucleotides or bases are at particular positions in at least two different specific polynucleotide sequences in material having a plurality of different polynucleotide sequences.

Background Art

At present, the most common technique for the detection of polynucleotide sequences is hybridisation using oligonucleotide or polynucleotide probes. For this technique, the target nucleic acid is usually bound, irreversibly, to a solid support such as cellulose or nylon. The labelled probe is then added in solution under hybridising conditions and if there is sequence homology between the probe and the target nucleic acid, the probe will form a hybrid molecule with the target polynucleotide. This hybrid can be detected in a variety of ways such as by radiolabelling or biotin labelling. The disadvantages of this technique are, firstly, that it requires considerable operator expertise, secondly, the technique is lengthy and time-consuming and thirdly, cannot be easily automated. Often the entire procedure can take more than 48 hours.

The liquid-solid methods normally employed for detecting specific nucleic acids in samples include Southern blot, Northern blot and dot blot hybridisations. These methods are slow, inefficient, technically demanding and are not easily automated. The Southern and Northern blot protocols have the further disadvantage of inefficient transfer of nucleic acid from gel to paper. Other groups have used liquid-solid hybridisations in which a capture probe is bound to a solid support and the DNA sequence of interest in the sample becomes bound to the capture probe. An example of

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this is in Australian Patent specification no. AU-70152/87 which describes using at least two oligonucleotide probes, which are complementary to mutually exclusive regions of the same target nucleic acid in a liquid hybridisation format to anneal the probes to the target, if it is present, and detecting the presence of the target by immobilisation of the two-probe target sandwich by one of the probes and subsequent detection of the other probe. However, this method requires a second, detector probe to hybridise to the DNA sequence of interest. This step reduces the specificity of the assay and subsequently increases background. The present invention overcomes this problem by involving only one probe which acts both as capture probe and as detector probe.

In contrast to liquid - solid hybridisation, liquid-liquid hybridisation has very rapid kinetics and improved sensitivity due to greater access of the probe to the target sequence. For example, Gene-Probe Inc uses a liquid hybridisation hydroxapatite method to detect DNA sequences. The main disadvantage of this system is that it relies on adsorptive discrimination of double-stranded DNA from single-stranded DNA sequences rather than sequence-specific separation of hybrid from excess probe. The present invention overcomes these disadvantages by allowing the nucleic acid hybridisations to occur in solution followed by the removal of the "hybrid" molecules onto a solid support matrix. Another potential advantage of liquid hybridisation is that a generalised solid support can work for a multitude of targets if the support - binding probes are labelled with the same capture molecule.

Several cases exist (Australian Patent specification nos. AU-A-70152/87, AU-A-26755/88, AU-A-53105/86, AU-B-89575/82 and AU-A-80097/87) which use a combination of two oligonucleotide probes to detect specific nucleic acid sequences in a sample. All these require the use of two short sequences of DNA on the target. These sequences must both be conserved in all possible target, must be mutually exclusive and non-overlapping and must have a similar G+C ratio to enable both probes to hybridise to their complementary sequence under the same conditions.

There is related background art concerning the use of a capture probe to detect the nucleic acid sequence of interest and to remove it from solution by binding the hybrid to a solid support matrix (Australian Patent specification nos. AU-A-70152/87, AU-A-53105/86, AU-A-21781/88, AU-A-69724/87, AU-A-14005/88). However, these techniques use separate

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capture and detector probes, resulting in a number of disadvantages as detailed above. The present invention overcomes these problems by using a single capture/detector probe system.

There are many examples available of attaching non-radioactive reporter molecules to DNA, to enable the detection of specific hybrids. However, when biotin is incorporated into the DNA molecule for detection, even though several biotin molecules may be incorporated per target molecule (thereby increasing the sensitivity of detection) the mechanism of visualising the incorporated biotin is complex and time consuming.

By contrast, the incorporation of other non-radioactive reporter molecules (such as fluorescent, luminescent, chemiluminescent molecules) enables rapid and simple detection of the target sequence. However, the present art only enables a single reporter molecule to be attached to each target sequence. This fact reduces the overall sensitivity of the final assay. The present invention overcomes both these problems at the one time by using a chemiluminescent detection system for simple, rapid detection and also by incorporating several detector molecules into each target, thereby significantly increasing assay sensitivity.

There is, therefore, a demand for a simple method which utilises the rapid kinetics of liquid hybridisation, which only requires a single probe for analysis, and which results in stable hybrids thereby allowing the unhybridised material to be easily removed from the sequences to be detected. Accordingly, the present invention provides a liquid hybridisation system in which a single probe hybridises to the sequence of interest and is then covalently extended to produce a stable hybrid. This hybrid is then captured on to a support matrix and subsequently washed to remove unhybridised material. The system described by the present invention is simple, rapid, sensitive; can be read visually or on a simple plate reader, and may be readily automated.

In this area of nucleic acid hybridisation there is a need to detect two broad types of diseases: infectious and genetic. In relation to infectious diseases, a number of DNA based systems have been described to detect diseases caused by bacteria such as *Salmonella*, *Neisseria*, parasitic organisms such as *Chlamydia* and *Rickettsiae*, viruses such as hepatitis B and protozoa such as *Plasmodium*. However, all of these suffer one or more of the disadvantages listed above.

In relation to genetic diseases which are characterised by a mutation

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(deletion, insertion, point mutation or translocation), the technology is less well developed. These types of diseases are currently diagnosed either by using restriction fragment length polymorphism (RFLP) analysis or by precise hybridisation of short oligonucleotide probes. RFLP detection requires that a restriction enzyme site is altered by the mutation and this is not always the case. In addition, RFLP analysis requires the use of Southern blot hybridisation for detection. The use of short oligonucleotide probes to detect point mutations also has several serious disadvantages. In particular, the hybridisation conditions must be precise to ensure that a single base-pair mismatch does not result in hybridisation. In practice, salt concentration and temperature determine the specificity of the hybridisation conditions and these are not easily controlled to the required preciseness. The present invention overcomes the need for Southern hybridisation analysis and for precise control of hybridisation conditions. It achieves this by the specific primer probe hybridising to a constant section of the gene adjacent to the mutation and allowing the enzyme, a DNA polymerase, to extend the DNA chain up to and including the nucleotide or base mutation. By manipulation of the dideoxynucleotide added to the polymerase reaction all of the possible nucleotide changes can be detected.

Detection of both infectious and genetic diseases requires the incorporation of some type of labelled molecule into the system. Various radioactive isotopes or radioactivity labelled compounds may be used as labels. However, radioactive labels have several disadvantages, including; (i) hazardous, (ii) expensive, (iii) limited shelf life, (iv) require expensive equipment for measuring the signal generated. More recently, a range of non-radioactive substances have been used to detect DNA molecules. Examples of non-radioactive labels are fluorescent, luminescent, chemiluminescent, enzymatic or immunological compounds. Labels based on the affinity of biotin and avidin or streptavidin, lanthanide chelates, lectins and proteins may also be used. The preferred detection means would be by spectroscopy or photochemistry or by the formation of a detectable complex between the label moiety and the polypeptide, lectin, or antibody linked to an entity capable of generating a detectable change, including enzymes such as; alkaline phosphatase or horseradish peroxidase.

However, what is lacking in the current technology is a single system

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which encompasses: (i) a rapid means of detecting a polynucleotide sequence, (ii) which is sufficiently sensitive to detect low numbers of the target sequence in the sample and (iii) which is non-radioactive. At present, no single system satisfies all these requirements.

As a final aspect, the need to detect specific polynucleotide sequences in a sample requires the organisation of all steps either; (i) into a simple kit format, or (ii) into an automated device. Whereas both, kits and automated machines, are available for detecting proteins by way of antibodies, no systems are yet available which simply, rapidly and inexpensively detect specific polynucleotide sequences in a sample.

Objects of Invention

It is an object of this invention to provide a method for detecting a specific polynucleotide sequence in a material.

Another object is to provide a method for detecting at least two different specific polynucleotide sequences in material having a plurality of different polynucleotide sequences.

A further object is to provide methods for detecting whether a specific nucleotide or base is at a particular position in a specific polynucleotide sequence.

Yet another object is to provide methods for detecting whether the same or different specific nucleotides or bases are at particular positions in at least two different specific polynucleotide sequences in material having a plurality of different polynucleotide sequences.

A further object is to provide a screening method for detecting the presence of at least two different specific polynucleotide sequences in material having a plurality of different polynucleotide sequences.

Another object is to provide screening methods for detecting whether the same or different specific nucleotides or bases are at particular positions in at least two different specific polynucleotide sequences in material having a plurality of different polynucleotide sequences.

Disclosure of Invention

The following abbreviations and definitions apply throughout the specification and claims:

- | | |
|--------|-----------------------------------------------------------------------------------------------------------------------|
| SP | - specific primer: a nucleotide sequence complementary to a portion of the target sequence |
| Target | - the nucleotide sequence to be detected in the sample; can be derived from, for example, any infectious or parasitic |

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organism including any virus, fungus, bacteria,
mycoplasma, nematode, protozoan etc

Polymerase	
Enzyme	- any DNA dependent DNA polymerase or RNA dependent DNA polymerase
dNTP	- all four deoxyribonucleotides that is dATP, dCTP, dGTP and dTTP as well as dITP and dUTP
ddNTP	- all four dideoxyribonucleotides that is ddATP, ddCTP, ddGTP and ddTTP as well as ddITP
D	- detector molecule: a molecule which can be covalently attached to a nucleotide or nucleotide sequence and that can be subsequently assayed for either directly or indirectly. For example, biotin; radioisotopes of carbon, hydrogen, iodine, phosphorus and sulphur; any antigen; haptens; fluorescent molecules including fluorescein, rhodamine and eosin; enzymes including alkaline phosphates, peroxidases and luciferase; any monoclonal antibody
dNTP-D	- a detector molecule covalently bound to one of the four deoxyribonucleotides
C	- capture molecule: a molecule which can be covalently attached to a nucleotide or nucleotide sequence and that will subsequently bind to a specific affinity molecule. For example, biotin (binds with avidin or streptavidin); antibody (binds with antigen); antigen (binds with antibody)
SAM	- specific affinity molecule: a molecule that will bind specifically with a particular capture molecule. For example, avidin or streptavidin; an antibody; an antigen
SS	- solid support: any solid matrix to which the SAM is attached. For example, agarose; polyacrylamide; magnetic beads; polystyrene; microtitre plates; nylon; nitrocellulose
Wash	- addition and removal of a solution for the purpose of removing unreacted molecules
Stringency	- conditions of salt and temperature which affect the stability of the hydrogen bonds between two nucleic acid

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molecules. For example, the hydrogen bonds are most unstable at high stringency reflecting either high temperature or low salt or both.

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|----------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Assay | <ul style="list-style-type: none"> - any procedure that is specific for detecting the detector molecule and that can be measured either qualitatively or quantitatively |
| X | <ul style="list-style-type: none"> - any one of the four deoxyribonucleotides |
| Y | <ul style="list-style-type: none"> - any one of the four ribonucleotides or deoxyribonucleotides; in the example Y will form hydrogen bonds with X, i.e., a nucleotide sequence of nine Y's will be complementary to a nucleotide sequence of 9 X's |
| Z | <ul style="list-style-type: none"> - any one of the four nucleotides forming a sequence not complementary to the sequence |
| ddT | <ul style="list-style-type: none"> - dideoxythymidine-5'-triphosphate: ddT has been used as example as it base pairs with A in the target sequence. If the base to be detected was C then ddG would be used, G with ddC and T with ddA. |
| ddT-C | <ul style="list-style-type: none"> - dideoxythymidine-5'-triphosphate/capture molecule complex: ddT has been used as example as it base pairs with A in the target sequence. If the base to be detected was C then ddG would be used, G with ddC and T with ddA. The capture molecule is covalently attached to ddT and represents any one of the capture molecules described above. |
| immediately adjacent | <ul style="list-style-type: none"> - means that there are no nucleotides or bases between a primer bound to part of a specific polynucleotide sequence and a specific nucleotide or base to be detected |
| fraction | <ul style="list-style-type: none"> - means at least a portion of a mixture resulting from the reaction of an oligonucleotide primer(s), extending reagents, specific polynucleotide sequence(s), detectable element(s) and/or separation elements |
| intervening sequence | <ul style="list-style-type: none"> - means at least one nucleotide or base between a primer bound to part of a specific polynucleotide sequence and a specific nucleotide or base to be detected |

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oligonucleotide

primer - a single stranded nucleic acid molecule with a typical length of 5 to 60 nucleotides but can be 200 or more nucleotides which has a sequence complementary to part of the polynucleotide sequence to be detected

specific

polynucleotide

sequence - a partly or completely known sequence of nucleotides

hybridisation - the physical association of the oligonucleotide primer with the complementary region of the target polynucleotide sequence to form a double stranded hybrid nucleic acid molecule

An interfering detectable and/or separation element is, for instance, one which is the same as that which is incorporated at the position complementary to the specific nucleotide or base to be detected, eg:

(A) Consider a single nucleotide or base N_1 to be detected in a specific polynucleotide sequence S_1 . Assume there is a bound oligonucleotide primer P_1 having a sequence complementary to part of S_1 and which is bound to S_1 . Assume there are intervening nucleotides N_A , N_B and N_C between P_1 and N_1 . Extend P_1 up to and including N_1 with complementary nucleotides to N_A , N_B , N_C and N_1 , namely, N_D , N_E , N_F and N_2-S_1 (S_1 corresponding to a separation element) respectively; then the following conditions are required:

N_A , N_B or N_C cannot be N_1

(B) Consider two nucleotides or bases N_1 and N_2 to be detected in two different specific polynucleotide sequences S_1 and S_2 . Assume there are two different oligonucleotide primers P_1 and P_2 (bound to S_1 and S_2 respectively) having sequence complementary to part of S_1 and S_2 .

Assume there are intervening nucleotides N_A , N_B and N_C between P_1 and N_1 and intervening nucleotides N_X , N_Y and N_Z between P_2 and N_2 . Extend (a) P_1 up to and including N_1 with complementary nucleotides to N_A , N_B , N_C and N_1 , namely, N_D , N_E , N_F and N_3-D_1 (D_1 corresponding to a detectable element) respectively; and (b) P_2 up to and including N_2 with complementary nucleotides to N_X , N_Y , N_Z and N_2 , namely, N_O , N_P , N_Q and N_4-D_2 (D_2

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corresponding to another detectable element);
then the following conditions are required:

If $N_1 = N_2$ and $D_1 = D_2$

then N_A)

N_B)

N_C) cannot be N_1

N_X)

N_Y)

N_Z)

If N_1 does not equal N_2

then N_A)

$D_1 = D_2$

N_B)

N_C) cannot be N_1 or N_2

N_X)

N_Y)

N_Z)

If N_1 does not equal N_2

N_A)

D_1 does not equal D_2

N_B)

N_C) cannot be N_1

N_X)

N_Y) cannot be N_2

N_Z)

According to a first embodiment of this invention there is provided a method for detecting a specific polynucleotide sequence in a material comprising:

- a) exposing said material to an oligonucleotide primer having a sequence complementary to part of said specific polynucleotide sequence wherein said primer binds to part of said polynucleotide sequence when present in said material;
- b) extending primer bound to the polynucleotide sequence wherein any extended primer includes a detectable element and/or a separation element;
- c) separating any extended primer into a fraction wherein said fraction does not have detectable element not included in said extended

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primer; and

- d) determining whether any extended primer is present in said fraction by assaying said fraction for said extended primer wherein the presence of said extended primer is indicative of the presence of the specific polynucleotide sequence in said material and the absence of said extended primer in said fraction is indicative of the absence of the specific polynucleotide sequence in said material.

According to a second embodiment of this invention there is provided a method for detecting at least two different specific polynucleotide sequences in material having a plurality of different polynucleotide sequences comprising:

- a) exposing said material to at least two different oligonucleotide primers each of said different oligonucleotide primers having a sequence complementary to part of one of said different specific polynucleotide sequences wherein each of said primers binds to part of one of said different specific polynucleotide sequences when present in said material each of said oligonucleotide primers binding to part of a different specific polynucleotide sequence to the other of said oligonucleotide primer(s);
- b) extending said different oligonucleotide primers bound to their different specific polynucleotide sequences wherein any extended primer includes a detectable element and/or a separation element;
- c) separating any extended primers into at least one fraction wherein said fraction(s) does not have detectable element not included in said extended primer(s); and
- d) determining whether any extended primers are present in said fraction(s) by assaying said fraction(s) for at least one of said extended primers wherein the presence of said extended primer(s) is indicative of the presence of at least one of the different specific polynucleotide sequences in said material and the absence of said extended primer(s) in said fraction(s) is indicative of the absence of at least one different specific polynucleotide sequence in said material.

According to a third embodiment of this invention there is provided a method for detecting whether a specific nucleotide or base is at a particular position in a specific polynucleotide sequence in material comprising:

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- a) exposing said material to an oligonucleotide primer having a sequence complementary to part of the specific polynucleotide sequence wherein said primer binds to part of said specific polynucleotide sequence in said material immediately adjacent to the particular position;
- b) extending primer bound to the specific polynucleotide sequence with the proviso that the bound primer is only extended up to and including said specific nucleotide or base when said specific nucleotide or base is at the particular position in the specific polynucleotide sequence wherein the extended primer has a detectable element and/or a separation element at a position complementary to said specific nucleotide or base;
- c) separating any extended primer into a fraction wherein said fraction does not have detectable element not included in said extended primer; and
- d) determining whether any extended primer is present in said fraction by assaying said fraction for said extended primer wherein the presence of said extended primer indicates that the specific nucleotide or base is at the particular position in the specific polynucleotide sequence and the absence of said extended primer indicates that the specific nucleotide or base is not at the particular position in the specific polynucleotide sequence.

According to a fourth embodiment of this invention there is provided a method for detecting whether the same or different specific nucleotides or bases are at particular positions in at least two different specific polynucleotide sequences in material having a plurality of different polynucleotide sequences comprising:

- a) exposing said material to at least two different oligonucleotide primers each of said different oligonucleotide primers having a sequence complementary to part of one of said different specific polynucleotide sequences wherein each of said primers binds to its complementary polynucleotide sequence when present in said material each of said oligonucleotide primers binding to part of a different specific polynucleotide sequence to that of the other primer(s) and wherein each of said primers binds to its complementary specific polynucleotide sequence in said material immediately adjacent to the particular position;
- b) extending said different oligonucleotide primers bound to their

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complementary polynucleotide sequences with the proviso that each of the bound primers is only extended up to and including the particular position when said specific nucleotide or base is at the particular position in the specific nucleotide sequence wherein each of said extended primers has a detectable element and/or a separation element at a position complementary to said specific nucleotide or base;

- c) separating any extended primer(s) which has extended only up to a specific nucleotide or base at a particular position into at least one fraction wherein said fraction does not have detectable element not included in said extended primer; and
- d) determining whether at least one extended primer is present in said fraction(s) by assaying said fraction(s) for said extended primer(s) wherein the presence of an extended primer indicates that a specific nucleotide or base is at a particular position in a specific polynucleotide sequence and the absence of said extended primer(s) in a fraction(s) indicates that at least one specific nucleotide or base is not at a particular position in at least one different specific polynucleotide sequence.

According to a fifth embodiment of this invention there is provided a method for detecting whether a specific nucleotide or base is at a particular position in a specific polynucleotide sequence in material comprising:

- a) exposing said material to an oligonucleotide primer having a sequence complementary to part of the specific polynucleotide sequence wherein said primer binds to part of said specific polynucleotide sequence in said material not immediately adjacent to the particular position whereby there is an intervening sequence between the particular position and primer bound to the specific polynucleotide sequence;
- b) extending bound primer with the proviso that the bound primer is only extended up to and including said specific nucleotide or base when said specific nucleotide or base is at the particular position in the specific polynucleotide sequence wherein the extended primer has a detectable element and/or a separation element at a position complementary to said specific nucleotide or base with the proviso that the intervening sequence cannot be one where bases or nucleotides complementary to the intervening sequence and which are incorporated into the extended primer cause incorporation of an

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- interfering detectable and/or separation element;
- c) separating any extended primer which has extended only up to a specific nucleotide or base at a particular position into a fraction wherein said fraction does not have detectable element not included in said extended primer; and
 - d) determining whether any extended primer is present in said fraction by assaying said fraction for said extended primer wherein the presence of said extended primer indicates that the specific nucleotide or base is at the particular position in the specific polynucleotide sequence and the absence of said extended primer indicates that the specific nucleotide or base is not at the particular position in the specific polynucleotide sequence.

According to a sixth embodiment of this invention there is provided a method for detecting whether the same or different specific nucleotides or bases are at particular positions in at least two different specific polynucleotide sequences in material having a plurality of different polynucleotide sequences comprising:

- a) exposing said material to at least two different oligonucleotide primers each of said different oligonucleotide primers having a sequence complementary to part of one of said different specific polynucleotide sequences wherein each of said primers binds to its complementary polynucleotide sequence not immediately adjacent to the particular position whereby there is an intervening sequence between the particular position and primer bound to the specific polynucleotide sequence when present in said material each of said oligonucleotide primers binding to part of a different specific polynucleotide sequence to that of the other primer(s);
- b) extending said different oligonucleotide primers bound to their complementary polynucleotide sequences wherein each of the bound primers is only extended up to and including the particular position when said specific nucleotide or base is at the particular position in the specific nucleotide sequence wherein each of said extended primers has a detectable element and/or a separation element at a position complementary to said specific nucleotide or base with the proviso that the intervening sequences cannot be ones where bases or nucleotides complementary to the intervening sequences and which are incorporated into the extended primer(s) cause incorporation of

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Interfering detectable and/or separation element(s);

- c) separating any extended primer which has extended only up to a specific nucleotide or base at a particular position into at least one fraction wherein said fraction(s) does not have detectable element not included in said extended primers; and
- d) determining whether any extended primers are present in said fraction(s) by assaying said fraction(s) for said extended primer(s) wherein the presence of an extended primer(s) indicates that at least one specific nucleotide or base is at a particular position in a specific polynucleotide sequence and the absence of said extended primer(s) in a fraction indicates that at least one specific nucleotide or base is not at a particular position in at least one different specific polynucleotide sequence.

According to a seventh embodiment of this invention there is provided a kit for detecting a specific polynucleotide sequence in a material comprising:

- a) an oligonucleotide primer having a sequence complementary to part of said specific polynucleotide sequence;
- b) polymerase enzyme for extended the oligonucleotide primer;
- c) at least one nucleotide selected from the group consisting of dATP, dCTP, dGTP, dTTP, dUTP, dITP, ddATP, ddCTP, ddGTP, ddITP and ddTTP; and
- d) a detectable element and/or separation element which is optionally attached to the oligonucleotide primer or one or more of the nucleotides.

According to an eighth embodiment of this invention there is provided a kit for detecting at least two different specific polynucleotide sequences in a material having a plurality of different polynucleotide sequences comprising:

- a) at least two different oligonucleotide primers each of said primers having a sequence complementary to part of one of said specific polynucleotide sequences;
- b) a polymerase enzyme(s) for extended the different oligonucleotide primers;
- c) at least one nucleotide selected from the group consisting of dATP, dCTP, dGTP, dTTP, dUTP, dITP, ddATP, ddCTP, ddGTP, ddITP and ddTTP; and

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- d) a detectable element and/or separation element which is optionally attached to the oligonucleotide primers or one or more of the nucleotides.

According to a ninth embodiment of this invention there is provided a kit for detecting whether a specific nucleotide or base is at a particular position in a specific polynucleotide sequence in a material comprising:

- a) an oligonucleotide primer having a sequence complementary to part of the specific polynucleotide sequence wherein said primer binds to part of said specific polynucleotide sequence immediately adjacent to the particular position;
- b) a polymerase enzyme for extending the oligonucleotide primer;
- c) one nucleotide selected from the group consisting of dATP, dCTP, dGTP, dTTP, dUTP, dITP, ddATP, ddCTP, ddGTP and ddTTP with the proviso that said nucleotide is selected whereby bound primer is only extended up to and including said specific nucleotide or base when said specific nucleotide or base is at the particular position in the specific polynucleotide sequence; and
- d) a detectable element and/or separation element whereby extended primer has a detectable element and/or a separation element at a position complementary to said specific nucleotide or base and a detectable and/or separation element optionally attached to the oligonucleotide primer.

According to a tenth embodiment of this invention there is provided a kit for detecting whether the same or different specific nucleotides or bases are at particular positions in at least two different specific polynucleotide sequences in material having a plurality of different polynucleotide sequences comprising:

- a) at least two different oligonucleotide primers each of said primers having a sequence complementary to part of one of said specific polynucleotide sequences wherein each of said primers binds to its complementary polynucleotide sequence when present in said material each of said oligonucleotide primers binding to part of a different specific polynucleotide sequence to that of the other primer(s) and wherein each of said primers binds to its complementary specific polynucleotide sequence in said material immediately adjacent to the particular position;
- b) a polymerase enzyme(s) for extending the different oligonucleotide

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primers;

- c) at least one nucleotide selected from the group consisting of dATP, dCTP, dGTP, dTTP, dUTP, dITP, ddITP, ddATP, ddCTP, ddGTP and ddTTP with the proviso that said nucleotide is selected whereby bound primer is only extended up to and including said specific nucleotide or base when said specific nucleotide or base is at the particular position in the specific polynucleotide sequence; and
- d) at least one detectable element and/or separation element whereby each extended primer has a detectable element and/or a separation element at a position complementary to said specific nucleotide or base and a detectable and/or separation element optionally attached to the oligonucleotide primer.

According to an eleventh embodiment of this invention there is provided a kit for detecting whether a specific nucleotide or base is at a particular position in a specific polynucleotide sequence in a material comprising:

- a) an oligonucleotide primer having a sequence complementary to part of the specific polynucleotide sequence wherein said primer binds to part of said specific polynucleotide sequence not immediately adjacent to the particular position whereby there is an intervening sequence between the particular position and primer bound to the specific polynucleotide sequence;
- b) a polymerase enzyme for extended the oligonucleotide primer;
- c) one nucleotide selected from the group consisting of dATP, dCTP, dGTP, dTTP, dUTP, dITP, ddITP, ddATP, ddCTP, ddGTP and ddTTP with the provision that said nucleotide is selected whereby bound primer is only extended up to and including said specific nucleotide or base when said specific nucleotide or base is at the particular position in the specific polynucleotide sequence with the proviso that the intervening sequence cannot be one where bases or nucleotides complementary to the intervening sequence and which are incorporated into the extended primer cause incorporation of an interfering detectable and/or separation element; and
- d) a detectable element and/or separation element whereby extended primer has a detectable element and/or a separation element at a position complementary to said specific nucleotide or base and a detectable and/or separation element optionally attached to the

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oligonucleotide primer.

According to a twelfth embodiment of this invention there is provided a kit for detecting whether the same or different specific nucleotides or bases are at particular positions in at least two different specific polynucleotide sequences in material having a plurality of different polynucleotide sequences comprising:

- a) at least two different oligonucleotide primers each of said primers having a sequence complementary to part of one of said specific polynucleotide sequences wherein each of said primers binds to its complementary polynucleotide sequence when present in said material each of said oligonucleotide primers binding to part of a different specific polynucleotide sequence to that of the other primer(s) and wherein each of said primers binds to its complementary specific polynucleotide sequence in said material not immediately adjacent to the particular position whereby there is an intervening sequence between the particular position and primer bound to the specific polynucleotide sequence;
- b) a polymerase enzyme(s) for extending the different oligonucleotide primers;
- c) at least one nucleotide selected from the group consisting of dATP, dCTP, dGTP, dTTP, dUTP, dITP, ddATP, ddCTP, ddGTP and ddTTP with the proviso that said nucleotide is selected whereby bound primer is only extended up to and including said specific nucleotide or base when said specific nucleotide or base is at the particular position in the specific polynucleotide sequence with the proviso that the intervening sequences cannot be ones where bases or nucleotides complementary to the intervening sequences and which are incorporated into the extended primer(s) cause incorporation of an interfering detectable and/or separation element(s); and
- d) at least one detectable element and/or separation element whereby each extended primer has a detectable element and/or a separation element at a position complementary to said specific nucleotide or base and a detectable and/or separation element optionally attached to the oligonucleotide primer.

According to a thirteenth embodiment of this invention there is provided an apparatus for performing the method of any one of the embodiments of the invention comprising:

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a reactor;

means for adding oligonucleotide primer and reagents for extending said primer to said reactor, said means for adding being operatively associated with the reactor;

means for separating extended primer into at least one fraction(s) and for holding said fraction(s), said means for separating being operatively associated with the reactor; and

a detector for detecting the presence of any extended primer(s) in said fraction(s), said detector being operatively associated with said means for separating.

According to a fourteenth embodiment of this invention there is provided a screening method for detecting the presence of at least two different specific polynucleotide sequences in material having a plurality of different polynucleotide sequences comprising:

- a) exposing said material to at least two different oligonucleotide primers each of said different oligonucleotide primers having a sequence complementary to part of one of said different specific polynucleotide sequences wherein each of said primers binds to part of one of said different specific polynucleotide sequences when present in said material each of said oligonucleotide primers binding to part of a different specific polynucleotide sequence to the other of said oligonucleotide primer(s);
- b) extending said different oligonucleotide primers bound to their different specific polynucleotide sequences wherein any extended primer includes a detectable element and/or a separation element;
- c) separating any extended primers into a fraction wherein said fraction does not have detectable element not included in said extended primer(s); and
- d) determining whether any extended primers are present in said fraction by assaying said fraction for all of said extended primers wherein the presence of any one of said extended primers is indicative of the present of at least one of the different specific polynucleotide sequences in said material and the absence of all of said extended primer(s) in said fraction is indicative of the absence of all of the different specific polynucleotide sequences in said material.

According to a fifteenth embodiment of this invention there is provided a screening method for detecting whether the same or different

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specific nucleotides or bases are at particular positions in at least two different specific polynucleotide sequences in material having a plurality of different polynucleotide sequences comprising:

- a) exposing said material to at least two different oligonucleotide primers each of said different oligonucleotide primers having a sequence complementary to part of one of said different specific polynucleotide sequences wherein each of said primers binds to its complementary polynucleotide sequence when present in said material each of said oligonucleotide primers binding to part of a different specific polynucleotide sequence to that of the other primer(s) and wherein each of said primers binds to its complementary specific polynucleotide sequence in said material immediately adjacent to the particular position;
- b) extending said different oligonucleotide primers bound to their complementary polynucleotide sequences with the proviso that each of the bound primers is only extended up to and including the particular position when said specific nucleotide or base is at the particular position in the specific nucleotide sequence wherein each of said extended primers has a detectable element and/or a separation element at a position complementary to said specific nucleotide or base;
- c) separating any extended primer(s) which has extended only up to a specific nucleotide or base at a particular position into a fraction wherein said fraction does not have detectable element not included in said extended primer; and
- d) determining whether at least one extended primer is present in said fraction by assaying said fraction for all of said extended primers wherein the present of an extended primer indicates that at least one specific nucleotide or base is at a particular position in a specific polynucleotide sequence and the absence of all said extended primers in said fraction indicates that at least one specific nucleotide or base is not at a particular position in any of the different specific polynucleotide sequences.

According to a sixteenth embodiment of this invention there is provided a screening method for detecting whether the same or different specific nucleotides or bases are at particular positions in at least two different specific polynucleotide sequences in material having a plurality of different polynucleotide sequences comprising:

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- a) exposing said material to at least two different oligonucleotide primers each of said different oligonucleotide primers having a sequence complementary to part of one of said different specific polynucleotide sequences wherein each of said primers binds to its complementary polynucleotide sequence not immediately adjacent to the particular position whereby there is an intervening sequence between the particular position and primer bound to the specific polynucleotide sequence when present in said material each of said oligonucleotide primers binding to part of a different specific polynucleotide sequence to that of the other primer(s);
- b) extending said different oligonucleotide primers bound to their complementary polynucleotide sequences wherein each of the bound primers is only extended up to and including the particular position when said specific nucleotide or base is at the particular position in the specific nucleotide sequence wherein each of said extended primers has a detectable element and/or a separation element at a position complementary to said specific nucleotide or base with the proviso that the intervening sequences cannot be ones where bases or nucleotides complementary to the intervening sequences and which are incorporated into the extended primer(s) cause incorporation of interfering detectable and/or separation element(s);
- c) separating any extended primer which has extended only up to a specific nucleotide or base at a particular position into a fraction wherein said fraction does not have detectable element not included in said extended primers; and
- d) determining whether any extended primers are present in said fraction by assaying said fraction for all of said extended primers wherein the presence of an extended primer indicates that at least one specific nucleotide or base is at a particular position in a specific polynucleotide sequence and the absence of any extended primer indicates that a specific nucleotide or base is not at a particular position in any of the different specific polynucleotide sequences. Generally the method for the detection of a nucleotide sequence or nucleotide change in a nucleotide sequence uses:
 - a nucleic acid primer specific to part of a partly or wholly known nucleotide sequence to be detected or specific to part of a partly or wholly known nucleotide sequence which is immediately adjacent to the

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nucleotide or base change to be detected or specific to part of a partly or wholly known nucleotide sequence which is not immediately adjacent to the nucleotide or base change to be detected but has an intervening sequence between the bound primer and the nucleotide or base change to be detected

extension of the primer catalysed by a nucleic acid polymerase enzyme either (i) the attachment of a capture molecule at the 5' end of the specific primer, addition of the target sequence under hybridisation conditions and the incorporation of a detector molecule(s) in the enzyme catalysed extended primer or (ii) the attachment of a detector molecule at the 5' end of the specific primer, addition of the target sequence under hybridisation conditions and the incorporation of a capture molecule(s) in the enzyme catalysed extended primer

capture of the extended primer using a specific affinity molecule attached to a solid support

assay for the detector molecule

Advantageously, for the first, second, seventh and eighth embodiments extending the primer comprises using at least one nucleotide selected from the group consisting of dATP, dCTP, dGTP, dTTP, dUTP, dITP, ddATP, ddCTP, ddGTP, ddITP and ddTTP. Typically two to four different nucleotides, and especially four different nucleotides are used.

Generally, for the third, fifth, ninth and eleventh embodiments extending the primer comprises using one nucleotide selected from the group consisting of dATP, dCTP, dGTP, dTTP, dUTP, dITP, ddATP, ddCTP, ddGTP, ddITP and ddTTP. Typically one nucleotide selected from the group consisting of ddATP, ddCTP, ddGTP, ddITP and ddTTP is used.

Typically, for the fourth, sixth, tenth and twelfth embodiments extending the primer comprises using at least one nucleotide selected from the group consisting of dATP, dCTP, dGTP, dTTP, dUTP, dITP, ddATP, ddCTP, ddGTP, ddITP and ddTTP. Generally at least one nucleotide selected from the group consisting of ddATP, ddCTP, ddGTP, ddITP and ddTTP is used.

In addition to a nucleotide(s) also typically used in extending the primer are at least one appropriate polymerase enzyme and appropriate buffering agent.

Examples of polymerase enzymes are E.coli DNA polymerase, E.coli DNA polymerase (Klenow fragment), Bacteriophage T7 DNA polymerase, Bacteriophage T4 DNA polymerase, Taq DNA polymerase and AMV Reverse

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transcriptase.

Buffering agents which buffer aqueous solutions between pH 6 - 8.5 are generally suitable for use with a polymerase enzyme to extend the primer. Generally a magnesium salt (eg MgCl₂ or MgSO₄) is included with the buffering agent. A total salt concentration between 50 and 300 mM is generally acceptable.

Temperature of the extending reaction is chosen according to the polymerase enzyme used and is typically in the range 25-80°C, more typically 30-70°C.

It is preferred that at least one of the nucleotides has the detectable element.

Alternatively, or in addition, the detectable element is linked to the oligonucleotide primer.

Typically, the detectable element is a radioactive label selected from the group consisting of ³H, ¹²⁵I, ¹³¹I, ¹⁴C, ³⁵S and ³²P.

The detectable element may be non-radioactive and generally is selected from the group consisting of an enzymatic group, an immunological group, and a spectroscopically detectable group. The spectroscopically detectable group may be a luminescent group, a chemiluminescent group, an NMR detectable group, a fluorescent group or an IR detectable group.

Advantageously, the extended primer includes a separation element which facilitates the separation of the extended primer into the first fraction.

Generally, the specific polynucleotide sequence is a DNA sequence or an RNA sequence.

The specific polynucleotide sequence may be from an infectious or disease causing organism which may be live or non-viable viral, bacterial, fungal or Protozoan.

The extending may include:

adding a plurality of nucleotide types to said primer bound to the polynucleotide sequence to extend said primer; or

adding a single nucleotide type to said primer bound to the polynucleotide sequence to extend said primer (third, fifth, ninth and eleventh embodiments); or

adding at least one single nucleotide type to said primer bound to the polynucleotide sequence to extend said primer.

In one particular method of the invention the exposure and primer

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extension steps occur before the separation step.

In one form of this invention, the separation element is linked to the oligonucleotide primer. Alternatively, the extended primer includes a separation element which facilitates separation of extended primer from the mixture into the first fraction and wherein at least one of said nucleotides has the separation element.

In one form of the invention, the separation step comprises:

contacting any extended primer with a molecule having affinity for the separation element, the molecule being linked to a support that facilitates the separating; and separating the contacted extended primer(s) to provide the fraction.

Advantageously, the separation element and the molecule are selected from the group listed immediately below:

Separation Element.

- (a) a ligand for which there is a receptor;
- (b) a receptor;
- (c) an antigen;
- (d) an antibody for an antigen;
- (e) an antiidiotypic antibody;
- (f) an antibody for an antiidiotypic antibody;
- (g) a haptenic group;
- (h) an antibody for a haptenic group;
- (i) an enzyme;
- (j) a binding inhibitor for an enzyme;

Molecule Having Affinity for Separation Element

- a receptor for the ligand;
- the ligand for the receptor;
- an antibody for the antigen;
- the antigen;
- an antibody for the antiidiotypic antibody;
- an antiidiotypic antibody for the antibody;
- an antibody for the haptenic group;
- the haptenic group;
- a binding inhibitor for the enzyme; and the enzyme.

Typically, the separation element and the molecule are selected from the group listed below:

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Separation Element

(a) a ligand for which there is a specific receptor;

(b) a specific receptor;

(c) an antigen;

(d) a specific antibody for an antigen;

(e) an antiidiotypic antibody;

(f) an antibody for an antiidiotypic antibody;

(g) a haptic group;

(h) a specific antibody for a haptic group;

(i) an enzyme;

(j) a tight binding inhibitor for an enzyme;

Typical examples of ligands for which there are available receptors are:

a vitamin such as biotin, a sugar molecule such as glucose or mannose, a hormone such as adrenalin or cortisone and a steroid such as progesterone or testosterone. There are numerous types of ligands which have available receptors and the preceding list is given by way of exemplification only.

The separation typically includes a solid support which is typically selected from the group consisting of latex, magnetic beads, nitrocellulose, agarose, cellulose, polyacrylamide, nylon, glass and polystyrene.

Advantages

1. General Advantages

A particular advantage of the method of the invention is that the system combines separation and detection elements on the one oligonucleotide after primer extension. Most other systems require the use of a capture probe and a detector probe.

A further advantage is that the system depends on the enzyme driven extension of the primer to incorporate a separation or detection element or

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elements (depending on configuration) and the element or elements are therefore covalently attached to the specific primer. Because of this covalent attachment, washing steps can be performed at very high stringency which results in reduced background and increased specificity.

Another advantage is that the method of the invention can be easily automated.

In the method of the invention the steps can be incorporated into a simple kit format.

A further advantage is that an oligonucleotide primer having a sequence complimentary to a conserved or variable region of the genome of the organism of interest can be used and thus can be designed for high or low specificity (i.e., genus specific, species specific or strain specific). a rapid and sensitive method for the detection of specific polynucleotide sequences or nucleotide changes in specific polynucleotide sequences using a single oligonucleotide probe.

Generally in the method of the invention the detectable hybrid is captured by a solid matrix to allow washing away of unincorporated detectable element. While this is not a difficult step it does add to the time required for processing. The step can be overcome by choosing a detectable element whereby any unincorporated detectable element can be simply inactivated and left in the reaction mix, rather than having to be physically removed.

Generally, a detection sensitivity in the order of $10^3 - 10^4$ genome copies is quite adequate for detecting a wide range of virus, bacterial and parasitic organisms which are usually present in moderate to high numbers in their respective disease states. For defect diseases in which the infectious agent is present only in very low numbers (e.g. HIV) the method of the invention can make use of the technique of polymerisation chain reaction (PCR) which is effectively an amplification step which results in an increase in sensitivity.

2. Infectious Disease Diagnosis Advantages

The method of the invention permits the rapid, simple and non-hazardous detection of specific polynucleotide sequences in samples of biological material especially infectious agents. The detectable sequence may be part of an infectious organism such as a virus, bacterium, Protozoan or fungus, or part of a gene in which an abnormality results in a genetic disorder: for example, sickle-cell anemia, cystic fibrosis.

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α -thalassemia or β -thalassemia.

Non-viable organisms can be detected using the method of the invention.

A further advantage is the method can use a mixture of oligonucleotide primers for the detection of a battery of agents suspected of causing a broad range of disease states (e.g., pneumonia; mycoplasma, chlamydia, streptococcus, legionella, RSV).

3. Genetic Disease Diagnosis Advantages

The method can be used for the rapid detection of an altered nucleotide base (point mutation) and for the detection of the insertion or deletion of one or more nucleotides in a polynucleotide sequence. In this instance the genetic change has to be known and characterised at the DNA sequence level.

The detection of a single or more base changes can be automated.

The method can be adapted to large scale screening for single (or more) nucleotide changes. This is particularly important in screening for genetic diseases such as cystic fibrosis but can also be adapted to the differentiation of alleles not necessarily involved in gene expression such as used for DNA profiling.

This particular method does not involve solid-liquid hybridisation. Other systems, such as Southern blot hybridisation, require that the target polynucleotide sequence is linked to a solid support (technically demanding) and/or the use of oligonucleotide probes that require precise hybridisation conditions. These systems cannot be automated and are not easily adapted to large scale screening.

Brief Description of Drawings

Preferred embodiments of the invention are now described with reference to the following drawings in which:

Figure 1 is a schematic drawing of a method for detecting a specific polynucleotide sequence;

Figure 2a is a schematic drawing of a method for detecting a specific polynucleotide sequence using a separation element which links to the oligonucleotide primer portion of an extended primer;

Figure 2b is a schematic drawing of a method for detecting a specific polynucleotide sequence using a separation element which links to the extended portion of an extended primer;

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Figure 3a is a schematic drawing of a method for detecting a nucleotide or base change using a separation element which links to the oligonucleotide primer portion of an extended primer;

Figure 3b is a schematic drawing of a method for detecting a nucleotide or base change using a separation element which links to the extended portion of an extended primer;

Figure 3c is a schematic drawing of a method for detecting a nucleotide or base change using a detectable element linked to the extended portion of an extended primer;

Figure 4 shows an autoradiograph used to verify the size of the extended oligonucleotide produced in the third example of the invention; and

Figure 5 is a schematic depiction of an apparatus for performing any one of the methods of the invention.

Best Mode and Other Modes of Carrying out the Invention

There are four especially preferred applications of the methods of the invention, that is, (i) the detection of a specific nucleotide sequence (major application: infectious diseases) (ii) the detection of a nucleotide or base change (major applications: genetic diseases and DNA fingerprinting) (iii) the detection of multiple specific nucleotide sequences using different capture molecules (major application: multiple infectious diseases) and (iv) detection of multiple nucleotide or base changes in different polynucleotide sequences (major applications: multiple genetic diseases and DNA fingerprinting). A single apparatus (with minor modifications) is suitable for use with all applications.

1. Detection of a specific nucleotide sequence

Referring to Figure 1, a method for detecting a specific polynucleotide sequence first involves the synthesis of an oligonucleotide primer complementary to the target polynucleotide sequence. The optimum length of the primer depends on the length of the most conserved sequence in the polynucleotide sequence to be detected up to a maximum length of 30 nucleotides.

The specific primer is preferably attached to an inert solid support, using the method of Ashley and MacDonald (1984). Any inert support particle to which small DNA molecules can be attached can be used, for instance cellulose or nylon. The polynucleotide sequence to be detected is added to the support - specific primer conjugate in the presence of a solution that promotes rapid hybridisation between the specific primer and

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the polynucleotide sequence. One appropriate solution contains 0.15M sodium chloride, 0.015M sodium citrate and 4% polyethylene glycol 6000. The support is then washed.

The choice of washing buffer determines the stringency conditions. A standard buffer is 140mM sodium chloride, 15mM sodium citrate and 20mM sodium phosphate, pH 7.0.

The specific primer is extended using either DNA polymerase I (Klenow fragment) or reverse transcriptase in the presence of the four deoxyribonucleotides, one of which is labelled, radioactively, with biotin or with a fluorescent label. The length of extension depends upon the time and temperature of the extension reaction providing all chemicals are in non-limiting concentration.

Generally the reaction is carried out at 37°C for 30 minutes and the label is ^{32}P . The level of incorporation of labelled nucleotide into the extended primer is measured by liquid scintillation when the label is ^{32}P .

Alternatively the label can be a fluorescent label.

Following primer extension the mixture is washed and assayed for presence of the label.

The presence of label after the final wash indicates that the primer has been extended, and that, therefore, the nucleic acid whose presence is being assayed is present. The specific primer can only be extended using the nucleic acid to which it is complementary as template.

In summary this method involves the following steps:

- the synthesis of a specific primer complementary to part of a known or partly known nucleic acid sequence, the target.

- the attachment of a capture (Configuration A - Figure 2a) or detector (Configuration B - Figure 2b) molecule to the 5' end of the specific primer

- the hybridisation, in solution, of this specific primer to the target nucleic acid sequence

- the addition to the solution of the four deoxyribonucleotides and either a DNA dependent DNA polymerase (if the target is DNA) or a RNA dependent DNA polymerase (if the target is RNA) under conditions suitable for the enzymatic extension of the specific primer. One or more of the deoxyribonucleotides has covalently attached to it a detector molecule (Configuration A) or a capture molecule

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(Configuration B)

the polymerase enzyme extends the primer using the target nucleic acid as a template and simultaneously incorporates either a detector or a capture molecule into the extended primer

the reaction can be heat denatured and the cycle repeated to amplify the amount of extended specific primer

for Configuration A, the specific affinity molecule (attached to a solid support) is added to the solution under conditions conducive to the binding of the capture molecule to the specific affinity molecule for Configuration B, the extended specific primer is precipitated with ethanol, washed to remove unincorporated deoxyribonucleotide-capture molecule complex. After washing, the extended specific primer is resuspended in a solution conducive to the binding of the capture molecule to the specific affinity molecule. The specific affinity molecule (attached to a solid support) is then added once the extended specific primer is attached to the specific affinity molecule-solid support complex via the capture molecule, the mix is washed extensively under high stringency conditions at the conclusion of the washing step, the mix is assayed using a procedure appropriate for detecting the detector molecule

2. Detection of a nucleotide or base change

This method involves the following steps:

the synthesis of a specific primer complementary to part of a known or partly known nucleic acid sequence, the target. The specific primer sequence is complementary to the target sequence either immediately adjacent to but not including the single nucleotide or base to be detected or not immediately adjacent to the single nucleotide or base to be detected but having an intervening sequence between the bound primer and the nucleotide or base to be detected the attachment of a capture (Configuration A - Figure 3a) or detector (Configuration B - Figure 3b) molecule to the 5' end of the specific primer. For Configuration C (Figure 3c), neither a capture or detector molecule are attached to the specific primer at the 5' end the hybridisation, in solution, of this specific primer to the target nucleic acid sequence

the addition to the solution of only one type of dideoxyribonucleotide (or deoxyribonucleotide), that is, the one which can pair

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or hydrogen bond with the base or nucleotide to be detected in the target sequence and either a DNA dependent DNA polymerase (if the target is DNA) or a RNA dependent DNA polymerase (if the target is RNA) under conditions suitable for the enzymatic extension of the specific primer. The dideoxyribonucleotide (or deoxyribo-nucleotide) has covalently attached to it a detector molecule (Configuration A and C) or a capture molecule (Configuration B) the polymerase enzyme extends the primer using the target nucleic acid as a template adding only one molecule of the dideoxyribonucleotide (or deoxyribonucleotide) if base pairing is possible. The specific primer is not extended if base pairing between the dideoxyribonucleotide (or deoxyribonucleotide) and the target sequence is not possible. A detector (Configuration A and C) or a capture (Configuration B) molecule

the reaction can be heat denatured and the cycle repeated to amplify the amount of extended specific primer

for Configuration A, the specific affinity molecule (attached to a solid support) is added to the solution under conditions conducive to the binding of the capture molecule to the specific affinity molecule for Configuration B, the extended specific primer is precipitated with ethanol, washed to remove unincorporated dideoxyribonucleotide-capture molecule complex (or deoxyribonucleotide-capture molecule complex). After washing, the extended specific primer is resuspended in a solution conducive to the binding of the capture molecule to the specific affinity molecule. The specific affinity molecule (attached to a solid support) is then added for Configurations A and B, once the extended specific primer is attached to the specific affinity molecule-solid support the complex via the capture molecule, the mix is washed extensively under high stringency conditions

for Configuration A and B, at the conclusion of the washing step, the mix is assayed using a procedure appropriate for detecting the detector molecule

for Configuration C, the reaction mix is loaded onto a lane of an agarose or polyacrylamide gel of suitable composition, electrophoresed to resolve the extended specific primer-detector molecule complex and finally assayed using a procedure appropriate

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for detecting the detector molecule. Labelled size standards are normally included in another lane of the gel.

3. Detection of Multiple Specific Nucleotide Sequences using Different Capture Molecules, i.e. Multiple Infectious Diseases

This method involves the following steps:

the synthesis of multiple different specific primers each of which is complementary to a different known nucleic acid sequence, the targets the attachment of different capture molecules to the 5' end of the different specific primers

the hybridisation, in solution, of these different specific primers (with attached different capture molecules) to the different target nucleic acid sequences to which the different specific primers are complementary

the addition to the solution of the four deoxyribonucleotides and either a DNA dependent DNA polymerase (if the targets are all DNA), or a RNA dependent DNA polymerase (if the targets are all RNA) or both polymerases (if the targets are a mixture of both DNA and RNA) under conditions suitable for the enzymatic extension of the specific primers if these specific primers are hybridised with their complementary target nucleic acid sequence. One or more of the deoxyribonucleotides has covalently attached to it a detector molecule the polymerase enzyme(s) extends the different specific primer using the different target nucleic acid sequences as a templates and simultaneously incorporates a detector molecule into the extended primers

the reaction can be heat denatured and the cycle repeated to amplify the amount of different extended specific primers

different specific affinity molecules (which have specific affinity for the different respective capture molecules and are attached to solid supports such as "test strips") are added to the solution under conditions conducive to the binding of the different capture molecule to their different specific affinity molecule

once the different extended specific primers are attached to their specific affinity molecule-solid support complexes via the different capture molecules, each different complex is removed individually with the appropriate test strip which subsequently is washed extensively under high stringency conditions

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at the conclusion of the washing step, one of the test strips is assayed for the presence of detector molecule using a procedure appropriate for the particular detector molecule. If the assay indicates that the detector molecule is present on the test strip then this is indicative that the specific nucleotide sequence targeted by the primer having that particular detectable molecule is present in the test sample. If the assay indicates that no detector molecule is present on the test strip then this is indicative that the specific nucleotide sequence targeted by the primer having that particular detectable molecule is not present in the test sample the procedure of the last step is repeated for each of the test strips

4. Detection of a multiple nucleotide or base changes in different polynucleotide sequences, ie multiple genetic defects or diseases or genetic fingerprinting

This method involves the following steps:

the synthesis of a multiple different specific primers each of which is complementary to a different known nucleic acid sequence, the targets. The different specific primer sequences are complementary to their respective different target sequences immediately adjacent to the single base to be detected

the attachment of different capture molecules to the 5' end of the different specific primers

the hybridization, in solution, of these different specific primers to their different respective target nucleic acid sequences

the addition of the solution of all four types of dideoxyribonucleotide (eg, ddATP, ddCTP, ddGTP, ddTTP) and either a DNA dependent DNA polymerase (if all the targets are DNA) or a RNA dependent DNA polymerase (if all the targets are RNA) or both (if the targets are a mixture of both DNA and RNA) under conditions suitable for the enzymatic extension of the different specific primers. Each dideoxyribonucleotide has covalently attached to it a different detector molecule.

the polymerase enzyme extends the different specific primers using the different target nucleic acid sequences as templates adding only one molecule of one of the dideoxyribonucleotides

the reaction can be heat denatured and the cycle repeated to amplify the amounts of extended different specific primers

the specific affinity molecules (with specific affinity for the different respective capture molecules and are attached to solid supports such as "test strips") are added to the solution under conditions conducive to the binding of the different capture molecules to the different specific affinity molecules once the different extended specific primers are attached to their specific affinity molecule-solid support complexes via the different capture molecules, each different complex is removed individually with the appropriate test strip which subsequently is washed extensively under high stringency conditions at the conclusion of the washing step, one of the test strips is assayed for the presence of detector molecule using a procedure appropriate for the particular detector molecule. If the assay indicates that the detector molecule is present on the test strip then this is indicative that the specific nucleotide sequence targeted by the primer having that particular detectable molecule is present in the test sample. If the assay indicates that no detector molecule is present on the test strip then this is indicative that the specific nucleotide sequence targeted by the primer having that particular detectable molecule is not present in the test sample the procedure of the last step is repeated for each of the test strips

5. Apparatus for Carrying out Methods of the Invention

Referring to Fig. 5 apparatus 100 is generally suitable for carrying out the methods of the invention according to non-automatic or automatic procedures. The extending and separation procedures take place within removable reactor 10 which is disposed within support vessel 11. Vessel 11 is temperature controlled by thermo-controlled jacket 12. The contents of reactor 10 are mixed by shaking vessel 11 using shaker 13. The base of reactor 10 has specific porosity membrane 14 supported by membrane support 15. Reactor 10 is supported in vessel 11 by support for removable reactor 21.

Sample material, primer(s) and extending reagents are delivered to reactor 10 from their respective vessels 16, 17 and 18. Primer annealing and extension are permitted to occur for an appropriate period under gentle mixing and fixed temperature conditions. This reaction creates an extended primer having a detectable element and separation element provided the sample has a specific polynucleotide sequence for which the primer has a

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sequence complementary to part of the specific polynucleotide sequence.

Subsequently specific affinity molecule(s) (SAM(S)) attached to an inert support for the separation element are added to reactor 10 from SAM supply vessel 19 and allowed to react with extended primer for an appropriate period. Unreacted reagents, buffer and unextended primer(s) are then removed to waste container 25 via vacuum line 25, having associated vacuum pump 24, attached to the base of support vessel 11 whilst extended primer(s) having attached SAM(S) are retained in reactor 10 by membrane 14. The retained primer-SAM complexes are washed with wash reagent supplied from vessel 22 to remove background and thereby create the first fraction in reactor 10. The first fraction is then examined for the presence of detectable element with detector 23. Typically, the detectable element is a radioactive element such as ^{32}P and detector 23 is a scintillation counter.

If the assay indicates that the detector molecule is present then this is indicative that the specific nucleotide sequence is present in the test sample. If the assay indicates that no detector molecule is present then this is indicative that the specific nucleotide sequence is not present in the test sample.

Example 1

In this example of the detection of a specific polynucleotide sequence, the method depicted in Fig. 1 was used for the detection of the single stranded genomic DNA of the bacteriophage M13.

1. Synthesis of the synthetic primer.

A 17-mer DNA complementary to M13 ssDNA was synthesized on an Applied Biosystems DNA Synthesizer.

The 17-mer had the nucleotide sequence:

3'-T-G-A-C-C-G-G-C-A-G-C-A-A-A-A-T-G- 5'

This primer is not only complementary to M13 single-stranded DNA but also to the bacterial double-stranded DNA plasmid, pUC12.

2. Attachment of the primer to cellulose.

The method of Ashley and MacDonald (1984) was used to attach the primer to ABM-cellulose. The amino benzyloxymethyl (ABM) groups on ABM cellulose were diazotized to form diazobenzyloxymethyl-cellulose (DBM-cellulose). The specific primer was attached to the DBM-cellulose. Approximately 40 μg of specific primer was bound to 20 mg of cellulose.

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3. Hybridization of specific primer-cellulose (SP-cellulose) with target nucleic acid

SP-cellulose was resuspended in a solution containing the target nucleic acid either M13 ssDNA or pUC12 and water. This was mixed, boiled for 5 minutes and quenched on ice. The mixture was then washed by centrifugation using a wash buffer comprising 140mM sodium chloride, 15mM sodium citrate and 20mM sodium phosphate, pH 7.0 to remove nucleic acids that had not hybridised with the SP-cellulose.

4. Primer extension

The specific primer was extended using DNA polymerase I (Klenow fragment) together with labelled and non-labelled nucleotides.

The reaction conditions were as follows:

20 : 1 Reaction Mix

50mM	Tris-HCl, pH 8.3
8mM	Magnesium chloride
4mM	dithiothreitol
4mM	sodium pyrophosphate
1mM	each dATP, dGTP and dTTP
1.CI	^{32}P dCTP

7 units DNA polymerase I, Klenow fragment.

It was found that the addition of polyethylene glycol 6000 to a final concentration of 4% increased the incorporation of radioactivity up to three fold.

The reaction mix was incubated for 2 hrs at 20°C..

At the end of the incubation, unincorporated nucleotides were removed by washing the SP-cellulose four times by centrifugation. The washing buffer was 140mM sodium chloride, 15mM sodium citrate and 20mM sodium phosphate, pH 7.0. The sample was vortexed for 1 minute at room temperature then centrifuged at 12,000g for 5 minutes. This was repeated four times.

Finally, the SP-cellulose was added to a vial containing scintillation solution and incorporation was determined using a liquid scintillation counter.

Example 2

In this example of the detection of a specific polynucleotide sequence according to the second embodiment of the invention, single stranded genomic DNA from the bacteriophage M13 mp18 was detected using the

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following oligonucleotide primer:

M13 ssDNA

3' . . . T T T G C T G C C G G T C A C G G T T C G A A . . . 5'

5' A A A C G A C G G C C A G T G C C

Specific Primer

The configuration used in this example is demonstrated in Figure 2b; the detector molecule in this example is ^{32}P and is attached to the 5' end of the specific primer; the capture molecule is biotin and is incorporated into the extended primer as biotin-16-dUTP (an analogue of dTTP); the specific affinity molecule is streptavidin and the solid support is agarose.

The specific primer was synthesised using an oligonucleotide synthesiser and was 5' end-labelled with ^{32}P using polynucleotide kinase as follows:

Reaction mix:

Specific primer (50 pmoles 5'OH ends)	1 μl
100 mM dithiothreitol	5 μl
10x TM Buffer (600 mM Tris, pH7.6; 90 mM MgCl ₂)	5 μl
γ - ^{32}P -dATP (3000 Ci/mmol)	15 μl
T ₄ polynucleotide kinase (8.6 units/ml)	2 μl
ddH ₂ O	22 μl

The reaction mix was incubated at 37°C for 20 minutes. The labelled primer was separated from unincorporated ^{32}P using a DEAE cellulose column.

The primer extension reaction was carried using the following reaction mix either with M13 mp18 or lambda bacteriophage (as a non-homologous control):

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	M13	Lambda
Template DNA (500 ng)	2.5ul	10ul
³² P labelled specific primer	10ul	10ul
DNA polymerase (Klenow; 1 unit/ul)	1ul	1ul
Biotin-16-dUTP (1mM)	1ul	1ul
dCTP (1 mM)	1ul	1ul
dGTP (1 mM)	1ul	1ul
dATP (1 mM)	1ul	1ul
NaCl (5 M)	1ul	1ul
MgSO ₄ (1 M)	0.5ul	0.5ul
Dithiothreitol (4 mM)	1.25ul	1.25ul
Bovine serum albumin (1 mg/ml)	2.5ul	2.5ul
Tris HCl, pH7.0 (1 M)	2.5ul	2.5ul
ddH ₂ O	24.75ul	25/25ul

The reaction mix was incubated at 37°C for 60 minutes. At the completion of the incubation, the extended primer was precipitated with ethanol to remove unincorporated biotin-16-dUTP. To the reaction mix was added one volume (50ul) 5M ammonium acetate and 250ul ethanol, followed by brief mixing and incubation at -20°C for one hour. The precipitate was collected by centrifugation, dried briefly, resuspended in 50ul ddH₂O and applied to a 200ul streptavidin/agarose column (containing 0.12mg streptavidin) previously equilibrated with binding buffer (10 mM Tris HCl, pH7.5; 200 mM NaCl; 1 mM EDTA). The column was washed five times with 500ul each of binding buffer. The streptavidin/agarose was suspended in 1ml binding buffer to which was added 6ml of scintillation fluid and the mixture counted in a liquid scintillation counter.

The results for the M13 and the lambda systems were as follows:

Template	CPM
M13	29,055
lambda	1,043

The results demonstrate that the primer was specifically extended only in the presence of the homologous template, M13, but not extended in the presence of the non-homologous template, lambda. Further, biotin was incorporated into the extended primer.

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Example 3

In this example of the use of the invention for the detection of an altered nucleotide base, the configuration shown in Figure 3c was used. The M13 sequence detected and the oligonucleotide primer sequence used were as follows:

M13 ssDNA
 3' . . . T T T G C T G C C G G T C A C G G T T C G A A . . . 5'
 5' A A A C G A C G G C C A G T G C C
 Specific Primer

The specific primer was synthesised using an oligonucleotide synthesiser.

The reaction mix was set up as follows:

500 ng M13mp18 ssDNA	2.5 μ l
0.8 pmoles specific primer	2.0 μ l
10x reaction buffer	1.5 μ l
ddH ₂ O	1.0 μ l

This reaction mix was incubated at 55°C for 10 minutes at which time the following were added:

³² P dATP (3000 Ci/mmol, 10 mCi/ml)	0.5 μ l
³² P ddATP (3000 Ci/mmol, 10 mCi/ml)	1.5 μ l
DNA polymerase (Klenow) (1 unit/ μ l)	1.0 μ l

This reaction mix was incubated for a further 15 minutes at 25°C. The reaction was stopped by the addition of 4 μ l formamide dye (95% formamide, 0.1% xylene cyanol, 0.1% bromophenol blue). The reaction mix was heated at 100°C for three minutes prior to loading on a 20% polyacrylamide gel and electrophoresing for 30 minutes at 30 milliamps. The gel was then fixed in 10% acetic acid, washed with ddH₂O and exposed to X-ray film at -80°C for 12 hours.

The resulting autoradiograph (Figure 4) exhibited only one band of 18 nucleotides in length. Thus the primer had been extended by only one nucleotide by the incorporation of a labelled adenine residue and thus detecting a specific single nucleotide in a nucleic acid sequence.

Example 4

In this example of the detection of a specific polynucleotide sequence, the method of Fig. 2a was used for detection of the single

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stranded genomic DNA of the bacteriophage M13.

1. Synthesis of the synthetic primer

A 25-mer DNA complementary to M13 ssDNA was synthesized by Clontec (USA).

The 25-mer had the nucleotide sequence

5'-X ACG TTG TAA AAC GAC GGC CAG TGC C-3'

where X is biotin covalently attached to a modified nucleotide.

2. Hybridisation and primer extension

The primer was mixed with buffer containing either M13 DNA (target) or herring sperm DNA (control) and extended using DNA Polymerase I (Klenow fragment).

Reaction conditions were as follows:

50 mM Tris pH 7.5

10 mM MgCl₂

4 mM DTT

100 mM NaCl

50 ug/ml bovine serum albumin

0.02 mM each dATP, dGTP, dTTP

1 Ci ³²P dCTP

742.5ng/50ul M13 or herring sperm DNA

25.8ng/50ul biotin-primer

0.5 units DNA polymerase I, Klenow fragment.

The reaction mixes were incubated at 37°C for 60' and terminated by precipitation and washing.

Precipitation was carried out by addition of 5M ammonium acetate, calf thymus DNA as carrier and ethanol. Pellets collected by centrifugation were washed with 70% ethanol until washings contained less than 500 cpm.

Washed pellets were then dissolved in binding buffer consisting of 200mM NaCl, 10mM Tris pH 7.5, 1mM EDTA and applied to a column of streptavidin-agarose. Columns were washed with several volumes of buffer and counts bound to streptavidin agarose (SA) were measured, with the following results:

Target	cpm bound to SA
M13	11,939
Herring Sperm	83

The results demonstrate that only primers extended in the presence of

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target M13 DNA result in significant radioactivity being bound to the column.

Industrial Applicability

The method of the invention can be readily used in the following applications:

- a) the simple, rapid, sensitive and automatable detection of infectious diseases of humans, animals and plants;
- b) the specific, rapid and large scale detection of base changes in nucleic acid sequences, particularly in the detection of gene defects and DNA fingerprinting;
- c) kits for use with the above applications; and
- d) automated equipment for use with the above applications.

CLAIMS

1. A method for detecting a specific polynucleotide sequence in a material comprising:
 - a) exposing said material to an oligonucleotide primer having a sequence complementary to part of said specific polynucleotide sequence wherein said primer binds to part of said polynucleotide sequence when present in said material;
 - b) extending primer bound to the polynucleotide sequence wherein any extended primer includes a detectable element or a separation element;
 - c) separating any extended primer into a fraction wherein said fraction does not have detectable element not included in said extended primer; and
 - d) determining whether any extended primer is present in said fraction by assaying said fraction for said extended primer wherein the presence of said extended primer is indicative of the presence of the specific polynucleotide sequence in said material and the absence of said extended primer in said fraction is indicative of the absence of the specific polynucleotide sequence in said material.
2. A method for detecting a specific polynucleotide sequence in a material comprising:
 - a) exposing said material to an oligonucleotide primer having a sequence complementary to part of said specific polynucleotide sequence wherein said primer binds to part of said polynucleotide sequence when present in said material;
 - b) extending primer bound to the polynucleotide sequence wherein any extended primer includes a detectable element and a separation element;
 - c) separating detectable element not included in said extended primer from any extended primer wherein any extended primer is separated into a fraction; and
 - d) determining whether any extended primer is present in said fraction by assaying said fraction for said detectable element wherein the presence of said detectable element is indicative of the presence of the specific polynucleotide sequence in said material and the absence of said detectable element in said fraction is indicative of the absence of the specific polynucleotide sequence in said material.

3. A method for detecting at least two different specific polynucleotide sequences in material having a plurality of different polynucleotide sequences comprising:

- a) exposing said material to at least two different oligonucleotide primers each of said different oligonucleotide primers having a sequence complementary to part of one of said different specific polynucleotide sequences wherein each of said primers binds to part of one of said different specific polynucleotide sequences when present in said material each of said oligonucleotide primers binding to part of a different specific polynucleotide sequence to the other of said oligonucleotide primer(s);
- b) extending said different oligonucleotide primers bound to their different specific polynucleotide sequences wherein any extended primer includes a detectable element or a separation element;
- c) separating any extended primers into at least one fraction wherein said fraction(s) does not have detectable element not included in said extended primer(s); and
- d) determining whether any extended primers are present in said fraction(s) by assaying said fraction(s) for at least one of said extended primers wherein the presence of said extended primer(s) is indicative of the presence of at least one of the different specific polynucleotide sequences in said material and the absence of said extended primer(s) in said fraction(s) is indicative of the absence of at least one different specific polynucleotide sequence in said material.

4. A method for detecting at least two different specific polynucleotide sequences in material having a plurality of different polynucleotide sequences comprising:

- a) exposing said material to at least two different oligonucleotide primers each of said different oligonucleotide primers having a sequence complementary to part of one of said different specific polynucleotide sequences wherein each of said primers binds to its complementary polynucleotide sequence when present in said material each of said oligonucleotide primers binding to part of a different specific polynucleotide sequence to that of the other oligonucleotide primers;

- b) extending said different oligonucleotide primers bound to their complementary polynucleotide sequences wherein any extended primer includes a detectable element and a separation element;
 - c) separating detectable element not included in said extended primers from any extended primers wherein any extended primers are separated into at least one fraction; and
 - d) determining whether any extended primers are present in said fraction(s) by assaying said fraction(s) for said detectable element(s) wherein the presence of said detectable element(s) is indicative of the presence of at least one of the different specific polynucleotide sequences in said material and absence of said detectable element(s) in a fraction(s) is indicative of the absence of at least one different specific polynucleotide sequence in said material.
5. A method for detecting whether a specific nucleotide or base is at a particular position in a specific polynucleotide sequence in material comprising:
- a) exposing said material to an oligonucleotide primer having a sequence complementary to part of the specific polynucleotide sequence wherein said primer binds to part of said specific polynucleotide sequence in said material immediately adjacent to the particular position;
 - b) extending primer bound to the specific polynucleotide sequence with the proviso that the bound primer is only extended up to and including said specific nucleotide or base when said specific nucleotide or base is at the particular position in the specific polynucleotide sequence wherein the extended primer has a detectable element and/or a separation element at a position complementary to said specific nucleotide or base;
 - c) separating any extended primer into a fraction wherein said fraction does not have detectable element not included in said extended primer; and
 - d) determining whether any extended primer is present in said fraction by assaying said fraction for said extended primer wherein the presence of said extended primer indicates that the specific nucleotide or base is at the particular position in the specific polynucleotide sequence and the absence of said extended primer indicates that the specific nucleotide or base is not at the particular position in the specific polynucleotide sequence.

6. A method for detecting whether the same or different specific nucleotides or bases are at particular positions in at least two different specific polynucleotide sequences in material having a plurality of different polynucleotide sequences comprising:
- a) exposing said material to at least two different oligonucleotide primers each of said different oligonucleotide primers having a sequence complementary to part of one of said different specific polynucleotide sequences wherein each of said primers binds to its complementary polynucleotide sequence when present in said material each of said oligonucleotide primers binding to part of a different specific polynucleotide sequence to that of the other primer(s) and wherein each of said primers binds to its complementary specific polynucleotide sequence in said material immediately adjacent to the particular position;
 - b) extending said different oligonucleotide primers bound to their complementary polynucleotide sequences with the proviso that each of the bound primers is only extended up to and including the particular position when said specific nucleotide or base is at the particular position in the specific nucleotide sequence wherein each of said extended primers has a detectable element and/or a separation element at a position complementary to said specific nucleotide or base;
 - c) separating any extended primer(s) which has extended only up to a specific nucleotide or base at a particular position into at least one fraction wherein said fraction does not have detectable element not included in said extended primer; and
 - d) determining whether at least one extended primer is present in said fraction(s) by assaying said fraction(s) for said extended primer(s) wherein the presence of an extended primer indicates that a specific nucleotide or base is at a particular position in a specific polynucleotide sequence and the absence of said extended primer(s) in a fraction(s) indicates that at least one specific nucleotide or base is not at a particular position in at least one different specific polynucleotide sequence.

7. A method for detecting whether a specific nucleotide or base is at a particular position in a specific polynucleotide sequence in material comprising:

- a) exposing said material to an oligonucleotide primer having a sequence complementary to part of the specific polynucleotide sequence wherein said primer binds to part of said specific polynucleotide sequence in said material not immediately adjacent to the particular position whereby there is an intervening sequence between the particular position and primer bound to the specific polynucleotide sequence;
- b) extending bound primer with the proviso that the bound primer is only extended up to and including said specific nucleotide or base when said specific nucleotide or base is at the particular position in the specific polynucleotide sequence wherein the extended primer has a detectable element and/or a separation element at a position complementary to said specific nucleotide or base with the proviso that the intervening sequence cannot be one where bases or nucleotides complementary to the intervening sequence and which are incorporated into the extended primer cause incorporation of an interfering detectable and/or separation element;
- c) separating any extended primer which has extended only up to a specific nucleotide or base at a particular position into a fraction wherein said fraction does not have detectable element not included in said extended primer; and
- d) determining whether any extended primer is present in said fraction by assaying said fraction for said extended primer wherein the presence of said extended primer indicates that the specific nucleotide or base is at the particular position in the specific polynucleotide sequence and the absence of said extended primer indicates that the specific nucleotide or base is not at the particular position in the specific polynucleotide sequence.

8. A method for detecting whether the same or different specific nucleotides or bases are at particular positions in at least two different specific polynucleotide sequences in material having a plurality of different polynucleotide sequences comprising:

- a) exposing said material to at least two different oligonucleotide primers each of said different oligonucleotide primers having a sequence complementary to part of one of said different specific polynucleotide sequences wherein each of said primers binds to its

complementary polynucleotide sequence not immediately adjacent to the particular position whereby there is an intervening sequence between the particular position and primer bound to the specific polynucleotide sequence when present in said material each of said oligonucleotide primers binding to part of a different specific polynucleotide sequence to that of the other primer(s);

- (b) extending said different oligonucleotide primers bound to their complementary polynucleotide sequences wherein each of the bound primers is only extended up to and including the particular position when said specific nucleotide or base is at the particular position in the specific nucleotide sequence wherein each of said extended primers has a detectable element and/or a separation element at a position complementary to said specific nucleotide or base with the proviso that the intervening sequences cannot be ones where bases or nucleotides complementary to the intervening sequences and which are incorporated into the extended primer(s) cause incorporation of interfering detectable and/or separation element(s);
- c) separating any extended primer which has extended only up to a specific nucleotide or base at a particular position into at least one fraction wherein said fraction(s) does not have detectable element not included in said extended primers; and
- d) determining whether any extended primers are present in said fraction(s) by assaying said fraction(s) for said extended primer(s)s wherein the presence of an extended primer(s) indicates that at least one specific nucleotide or base is at a particular position in a specific polynucleotide sequence and the absence of said extended primer(s) in a fraction indicates that at least one specific nucleotide or base is not at particular positions in at least one different specific polynucleotide sequence.

9. The method of claim 1 wherein said extending comprises using at least one nucleotide selected from the group consisting of dATP, dCTP, dGTP, dTTP, dUTP, dITP, ddATP, ddCTP, ddGTP, ddITP and ddTTP.

10. The method of claim 2 wherein said extending comprises using at least one nucleotide selected from the group consisting of dATP, dCTP, dGTP, dTTP, dUTP, dITP, ddATP, ddCTP, ddGTP, ddITP and ddTTP.

11. The method of claim 3 wherein said extending comprises using at least one nucleotide selected from the group consisting of dATP, dCTP, dGTP, dTTP, dUTP, dITP, ddATP, ddCTP, ddGTP, ddITP and ddTTP.
 12. The method of claim 4 wherein said extending comprises using at least one nucleotide selected from the group consisting of dATP, dCTP, dGTP, dTTP, dUTP, dITP, ddATP, ddCTP, ddGTP, ddITP and ddTTP.
 13. The method of claim 1 wherein said extending comprises using four different nucleotides selected from the group consisting of dATP, dCTP, dGTP, dTTP, dUTP, dITP, ddATP, ddCTP, ddGTP, ddITP and ddTTP.
 14. The method of claim 2 wherein said extending comprises using four different nucleotides selected from the group consisting of dATP, dCTP, dGTP, dTTP, dUTP, dITP, ddATP, ddCTP, ddGTP, ddITP and ddTTP.
 15. The method of claim 3 wherein said extending comprises using four different nucleotides selected from the group consisting of dATP, dCTP, dGTP, dTTP, dUTP, dITP, ddATP, ddCTP, ddGTP, ddITP and ddTTP.
 16. The method of claim 4 wherein said extending comprises using four different nucleotides selected from the group consisting of dATP, dCTP, dGTP, dTTP, dUTP, dITP, ddATP, ddCTP, ddGTP, ddITP and ddTTP.
 17. The method of claim 5 wherein said extending comprises using one nucleotide selected from the group consisting of dATP, dCTP, dGTP, dTTP, dUTP, dITP, ddITP, ddATP, ddCTP, ddGTP and ddTTP.
 18. The method of claim 5 wherein said extending comprises using one nucleotide selected from the group consisting of ddITP, ddATP, ddCTP, ddGTP and ddTTP.
 19. The method of claim 6 wherein said extending comprises using at least one nucleotide selected from the group consisting of dATP, dCTP, dGTP, dTTP, dUTP, dITP, ddATP, ddCTP, ddGTP, ddITP and ddTTP.
 20. The method of claim 6 wherein said extending comprises using at least one nucleotide selected from the group consisting of ddATP, ddCTP, ddGTP, ddITP and ddTTP.
 21. The method of claim 7 wherein said extending comprises using one nucleotide selected from the group consisting of dATP, dCTP, dGTP, dTTP, dUTP, dITP, ddITP, ddATP, ddCTP, ddGTP and ddTTP.

22. The method of claim 8 wherein said extending comprises using at least one nucleotide selected from the group consisting of dATP, dCTP, dGTP, dTTP, dUTP, dITP, ddATP, ddCTP, ddGTP, ddITP and ddTTP.
23. The method of any one of claims 9 to 22 wherein at least one of said nucleotides has a detectable element.
24. The method of any one of claims 9 to 22 wherein at least one of said nucleotides has a separation element.
25. The method of any one of claims 9 to 22 wherein at least one of said nucleotides has separation and detectable elements.
26. The method of any one of claims 1 to 8 wherein the oligonucleotide primer incorporates a detectable element.
27. The method of any one of claims 1 to 8 wherein the oligonucleotide primer incorporates a separation element.
28. The method of any one of claims 1 to 8 wherein the oligonucleotide primer incorporates detectable and separation elements.
29. The method of any one of claims 1 to 8 wherein any extended primer includes a separation element and said separating step comprises: contacting any extended primer with a molecule having affinity for the separation element, said molecule being linked to a support that facilitates said separating; and separating said contacted extended primer(s) to provide said fraction.
30. The method of any one of claims 1 to 8 wherein any extended primer includes a separation element and said separating step comprises: contacting any extended primer with a molecule having affinity for the separation element; and separating said contacted extended primer(s) to provide said fraction wherein said separation element and said molecule are selected from the group listed as follows:

	<u>Separation Element</u>	<u>Molecule Having Affinity for Separation Element</u>
(a)	a ligand for which there is a specific receptor	a specific receptor for the ligand;
(b)	a specific receptor;	the ligand for the specific receptor;
(c)	an antigen;	a specific antibody for the antigen;
(d)	a specific antibody for an antigen;	the antigen;
(e)	an antiidiotypic antibody	a specific antibody for the antiidiotypic antibody;
(f)	an antibody for an antiidiotypic antibody;	an antiidiotypic antibody specific for the antibody
(g)	a haptenic group;	a specific antibody for the haptenic group;
(h)	a specific antibody for a haptenic group;	the haptenic group
(i)	an enzyme;	a tight binding inhibitor for the enzyme; and
(j)	a tight binding inhibitor for an enzyme;	the enzyme.
32.	The method of any one of claims 1 to 8 wherein the specific polynucleotide sequence(s) is a DNA sequence.	
33.	The method of any one of claims 1 to 8 wherein the specific polynucleotide sequence(s) is an RNA sequence.	
34.	The method of any one of claims 1 to 4 wherein the specific polynucleotide sequence(s) is from an infectious or disease causing organism.	
35.	A kit for detecting a specific polynucleotide sequence in a material comprising:	
a)	an oligonucleotide primer having a sequence complementary to part of said specific polynucleotide sequence;	
b)	polymerase enzyme for extending the oligonucleotide primer;	

Separation Element

- (a) a ligand for which there is a receptor
- (b) a receptor;
- (c) an antigen;
- (d) an antibody for an antigen
- (e) an antiidiotypic antibody
- (f) an antibody for an antiidiotypic antibody;
- (g) a haptic group;
- (h) an antibody for a haptic group;
- (i) an enzyme;
- (j) a binding inhibitor for an enzyme;

31.

The method of any one of claims 1 to 6 wherein any extended primer includes a separation element and said separating step comprises: contacting any extended primer with a molecule having affinity for the separation element, said molecule being linked to a support that facilitates said separating; and separating said contacted extended primer(s) to provide said fraction wherein said separation element and said molecule are selected from the group listed as follows:

Molecule Having Affinity for Separation Element

- a receptor for the ligand;
- the ligand for the receptor;
- an antibody for the antigen;
- the antigen;
- an antibody for the antiidiotypic antibody;
- an antiidiotypic antibody for the antibody
- an antibody for the haptic group;
- the haptic group
- a binding inhibitor for the enzyme; and
- the enzyme.

d) a detectable element and/or separation element whereby extended primer has a detectable element and/or a separation element at a position complementary to said specific nucleotide or base and a detectable and/or separation element optionally attached to the oligonucleotide primer.

38. A kit for detecting whether the same or different specific nucleotides or bases are at particular positions in at least two different specific polynucleotide sequences in material having a plurality of different polynucleotide sequences comprising:

a) at least two different oligonucleotide primers each of said primers having a sequence complementary to part of one of said specific polynucleotide sequences wherein each of said primers binds to its complementary polynucleotide sequence when present in said material each of said oligonucleotide primers binding to part of a different specific polynucleotide sequence to that of the other primer(s) and wherein each of said primers binds to its complementary specific polynucleotide sequence in said material immediately adjacent to the particular position;

b) a polymerase enzyme(s) for extending the different oligonucleotide primers;

c) at least one nucleotide selected from the group consisting of dATP, dCTP, dGTP, dTTP, dUTP, dITP, ddITP, ddATP, ddCTP, ddGTP and ddTTP with the proviso that said nucleotide is selected whereby bound primer is only extended up to and including said specific nucleotide or base when said specific nucleotide or base is at the particular position in the specific polynucleotide sequence; and

d) at least one detectable element and/or separation element whereby each extended primer has a detectable element and/or a separation element at a position complementary to said specific nucleotide or base and a detectable and/or separation element optionally attached to the oligonucleotide primer.

39. A kit for detecting whether a specific nucleotide or base is at a particular position in a specific polynucleotide sequence in a material comprising:

a) an oligonucleotide primer having a sequence complementary to part of the specific polynucleotide sequence wherein said primer binds to part of said specific polynucleotide sequence not immediately adjacent to the particular

c) at least one nucleotide selected from the group consisting of dATP, dCTP, dGTP, dTTP, dUTP, dITP, ddATP, ddCTP, ddGTP, ddITP and ddTTP; and

d) a detectable element and/or separation element which is optionally attached to the oligonucleotide primer or one or more of the nucleotides.

36. A kit for detecting at least two different specific polynucleotide sequences in a material having a plurality of different polynucleotide sequences comprising:

a) at least two different oligonucleotide primers each of said primers having a sequence complementary to part of one of said specific polynucleotide sequences;

b) a polymerase enzyme(s) for extending the different oligonucleotide primers;

c) at least one nucleotide selected from the group consisting of dATP, dCTP, dGTP, dTTP, dUTP, dITP, ddATP, ddCTP, ddGTP, ddITP and ddTTP; and

d) a detectable element and/or separation element which is optionally attached to the oligonucleotide primers or one or more of the nucleotides.

37. A kit for detecting whether a specific nucleotide or base is at a particular position in a specific polynucleotide sequence in a material comprising:

a) an oligonucleotide primer having a sequence complementary to part of the specific polynucleotide sequence wherein said primer binds to part of said specific polynucleotide sequence immediately adjacent to the particular position;

b) a polymerase enzyme for extending the oligonucleotide primer;

c) one nucleotide selected from the group consisting of dATP, dCTP, dGTP, dTTP, dUTP, dITP, ddATP, ddCTP, ddGTP and ddTTP with the proviso that said nucleotide is selected whereby bound primer is only extended up to and including said specific nucleotide or base when said specific nucleotide or base is at the particular position in the specific polynucleotide sequence; and

WU WU/WY4CC

position whereby there is an intervening sequence between the particular position and primer bound to the specific polynucleotide sequence;

- b) a polymerase enzyme for extending the oligonucleotide primer;
- c) one nucleotide selected from the group consisting of dATP, dCTP, dGTP, dTTP, dUTP, dITP, ddATP, ddCTP, ddGTP and ddTTP with the proviso that said nucleotide is selected whereby bound primer is only extended up to and including said specific nucleotide or base when said specific nucleotide or base is at the particular position in the specific polynucleotide sequence with the proviso that the intervening sequence cannot be one where bases or nucleotides complementary to the intervening sequence and which are incorporated into the extended primer cause incorporation of an interfering detectable and/or separation element; and
- d) a detectable element and/or separation element whereby extended primer has a detectable element and/or a separation element at a position complementary to said specific nucleotide or base and a detectable and/or separation element optionally attached to the oligonucleotide primer.

40. A kit for detecting whether the same or different specific nucleotides or bases are at particular positions in at least two different specific polynucleotide sequences in material having a plurality of different polynucleotide sequences comprising:

- a) at least two different oligonucleotide primers each of said primers having a sequence complementary to part of one of said specific polynucleotide sequences wherein each of said primers binds to its complementary polynucleotide sequence when present in said material each of said oligonucleotide primers binding to part of a different specific polynucleotide sequence to that of the other primer(s) and wherein each of said primers binds to its complementary specific polynucleotide sequence in said material not immediately adjacent to the particular position whereby there is an intervening sequence between the particular position and primer bound to the specific polynucleotide sequence;
- b) a polymerase enzyme(s) for extending the different oligonucleotide primers;
- c) at least one nucleotide selected from the group consisting of dATP, dCTP, dGTP, dTTP, dUTP, dITP, ddITP, ddATP, ddCTP, ddGTP and ddTTP with the proviso that said

nucleotide is selected whereby bound primer is only extended up to and including said specific nucleotide or base when said specific nucleotide or base is at the particular position in the specific polynucleotide sequence with the proviso that the intervening sequences cannot be ones where bases or nucleotides complementary to the intervening sequences and which are incorporated into the extended primer(s) cause incorporation of an interfering detectable and/or separation element(s); and

- d) at least one detectable element and/or separation element whereby each extended primer has a detectable element and/or a separation element at a position complementary to said specific nucleotide or base and a detectable and/or separation element optionally attached to the oligonucleotide primer.

41. An apparatus for performing the method of any one of claims 1 to 8 comprising:

a reactor;

means for adding oligonucleotide primer and reagents for extending said primer to said reactor, said means for adding being operatively associated with the reactor;

means for separating extended primer into at least one fraction(s) and for holding said fraction(s), said means for separating being operatively associated with the reactor; and

a detector for detecting the presence of any extended primer(s) in said fraction(s), said detector being operatively associated with said means for separating.

42. A screening method for detecting the presence of at least two different specific polynucleotide sequences in material having a plurality of different polynucleotide sequences comprising:
- exposing said material to at least two different oligonucleotide primers each of said different oligonucleotide primers having a sequence complementary to part of one of said different specific polynucleotide sequences wherein each of said primers binds to part of one of said different specific polynucleotide sequences when present in said material each of said oligonucleotide primers binding to part of a different specific polynucleotide sequence to the other of said oligonucleotide primer(s);
 - extending said different oligonucleotide primers bound to their different specific polynucleotide sequences wherein any extended primer includes a detectable element and/or a separation element;
 - separating any extended primers into a fraction wherein said fraction does not have detectable element not included in said extended primer(s); and
 - determining whether any extended primers are present in said fraction by assaying said fraction for all of said extended primers wherein the presence of any one of said extended primers is indicative of the present of at least one of the different specific polynucleotide sequences in said material and the absence of all of said extended primer(s) in said fraction is indicative of the absence of all of the different specific polynucleotide sequences in said material.
43. A screening method for detecting whether the same or different specific nucleotides or bases are at particular positions in at least two different specific polynucleotide sequences in material having a plurality of different polynucleotide sequences comprising:
- exposing said material to at least two different oligonucleotide primers each of said different oligonucleotide primers having a sequence complementary to part of one of said different specific polynucleotide sequences wherein each of said primers binds to its complementary polynucleotide sequence when present in said material each of said oligonucleotide primers binding to part of a different specific polynucleotide sequence to that of the other primer(s) and wherein each of said primers binds to its complementary specific

- polynucleotide sequence in said material immediately adjacent to the particular position;
- b) extending said different oligonucleotide primers bound to their complementary polynucleotide sequences with the proviso that each of the bound primers is only extended up to and including the particular position when said specific nucleotide or base is at the particular position in the specific nucleotide sequence wherein each of said extended primers has a detectable element and/or a separation element at a position complementary to said specific nucleotide or base;
- c) separating any extended primer(s) which has extended only up to a specific nucleotide or base at a particular position into a fraction wherein said fraction does not have detectable element not included in said extended primer; and
- d) determining whether at least one extended primer is present in said fraction by assaying said fraction for all of said extended primers wherein the present of an extended primer indicates that at least one specific nucleotide or base is at a particular position in a specific polynucleotide sequence and the absence of all said extended primers in said fraction indicates that at least one specific nucleotide or base is not at a particular position in any of the different specific polynucleotide sequences.
44. A screening method for detecting whether the same or different specific nucleotides or bases are at particular positions in at least two different specific polynucleotide sequences in material having a plurality of different polynucleotide sequences comprising:
- a) exposing said material to at least two different oligonucleotide primers each of said different oligonucleotide primers having a sequence complementary to part of one of said different specific polynucleotide sequences wherein each of said primers binds to its complementary polynucleotide sequence not immediately adjacent to the particular position whereby there is an intervening sequence between the particular position and primer bound to the specific polynucleotide sequence when present in said material each of said oligonucleotide primers binding to part of a different specific polynucleotide sequence to that of the other primer(s);
- b) extending said different oligonucleotide primers bound to their

complementary polynucleotide sequences wherein each of the bound primers is only extended up to and including the particular position when said specific nucleotide or base is at the particular position in the specific nucleotide sequence wherein each of said extended primers has a detectable element and/or a separation element at a position complementary to said specific nucleotide or base with the proviso that the intervening sequences cannot be ones where bases or nucleotides complementary to the intervening sequences and which are incorporated into the extended primer(s) cause incorporation of interfering detectable and/or separation element(s);

- c) separating any extended primer which has extended only up to a specific nucleotide or base at a particular position into a fraction wherein said fraction does not have detectable element not included in said extended primers; and
- d) determining whether any extended primers are present in said fraction by assaying said fraction for all of said extended primers wherein the presence of an extended primer indicates that at least one specific nucleotide or base is at a particular position in a specific polynucleotide sequence and the absence of any extended primer indicates that a specific nucleotide or base is not at a particular position in any of the different specific polynucleotide sequences.

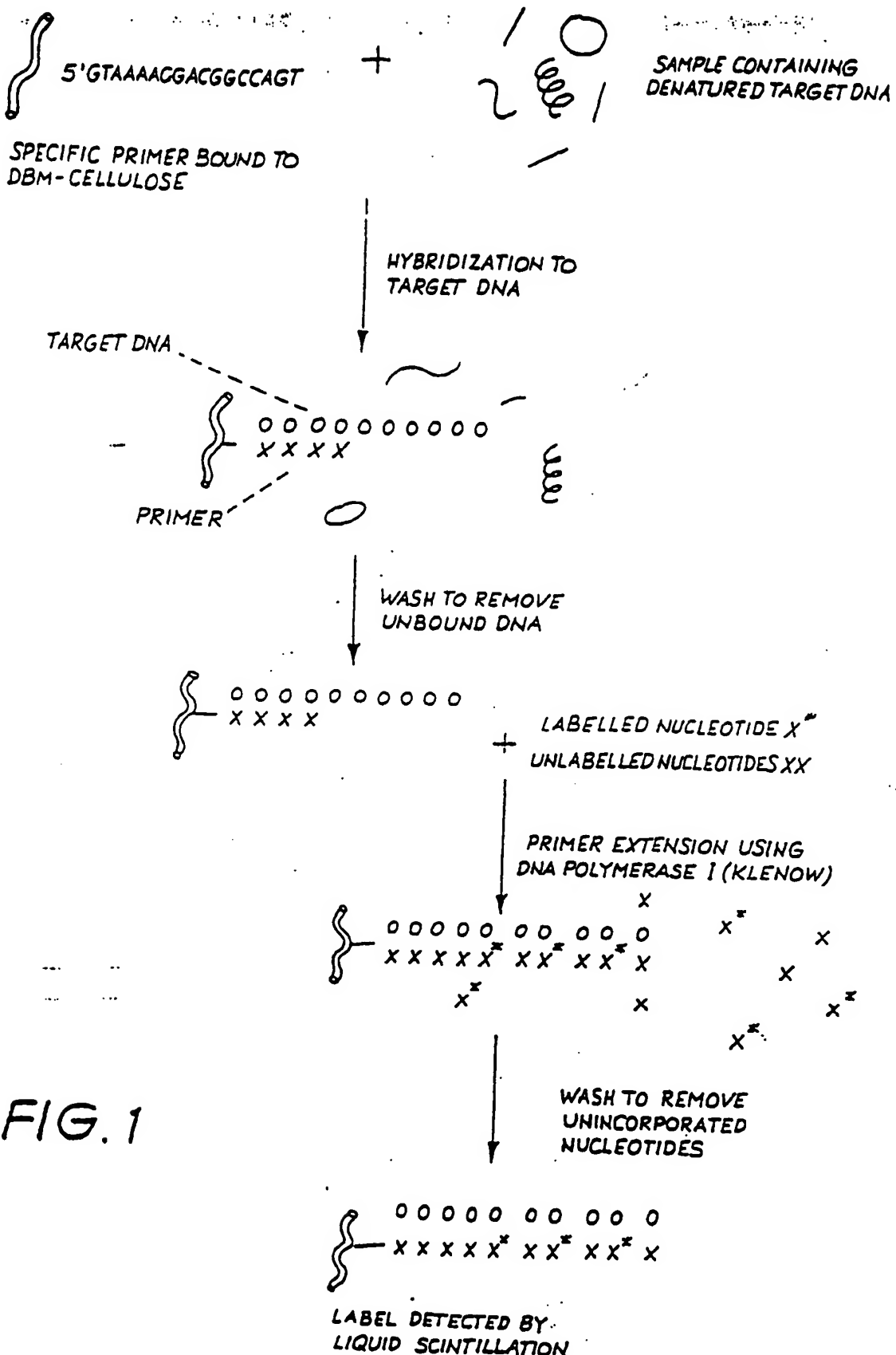


FIG. 1

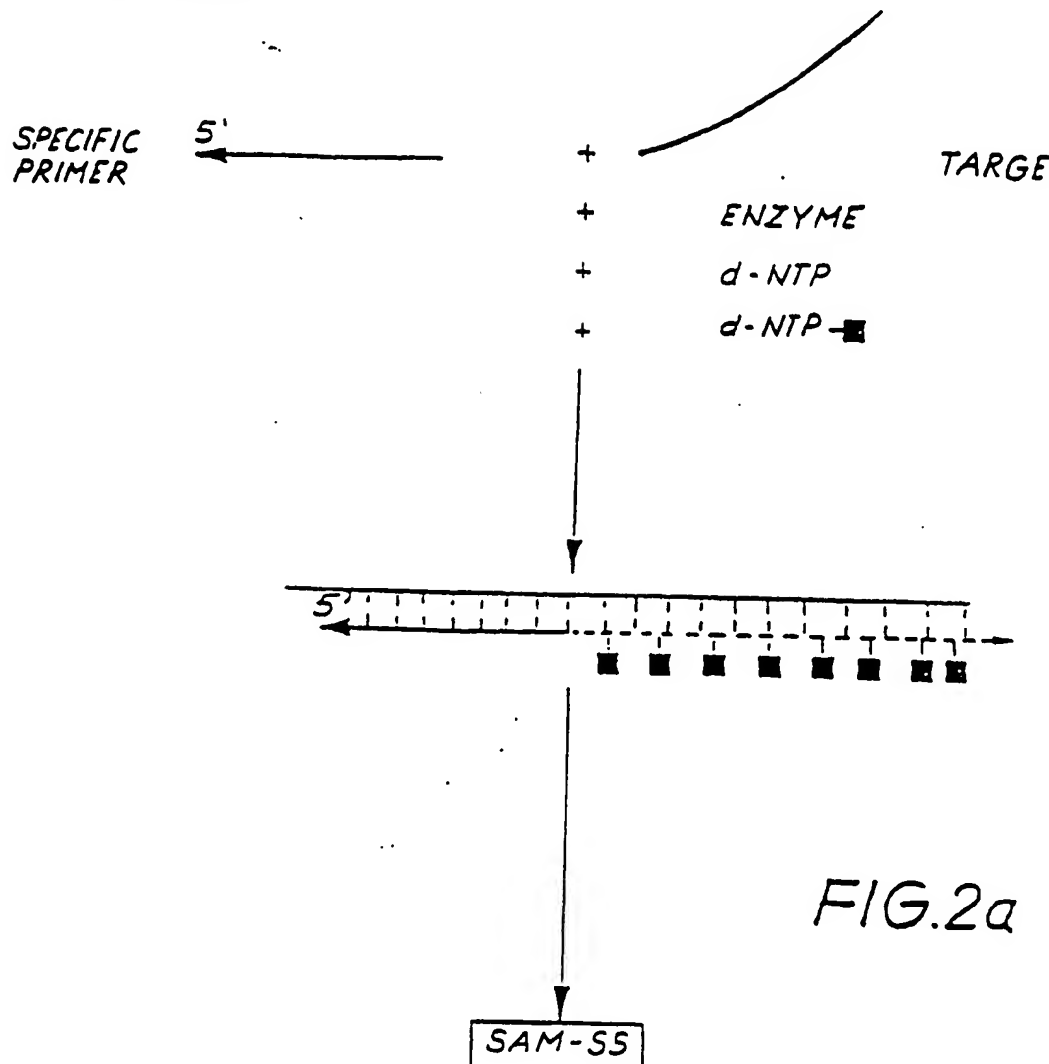
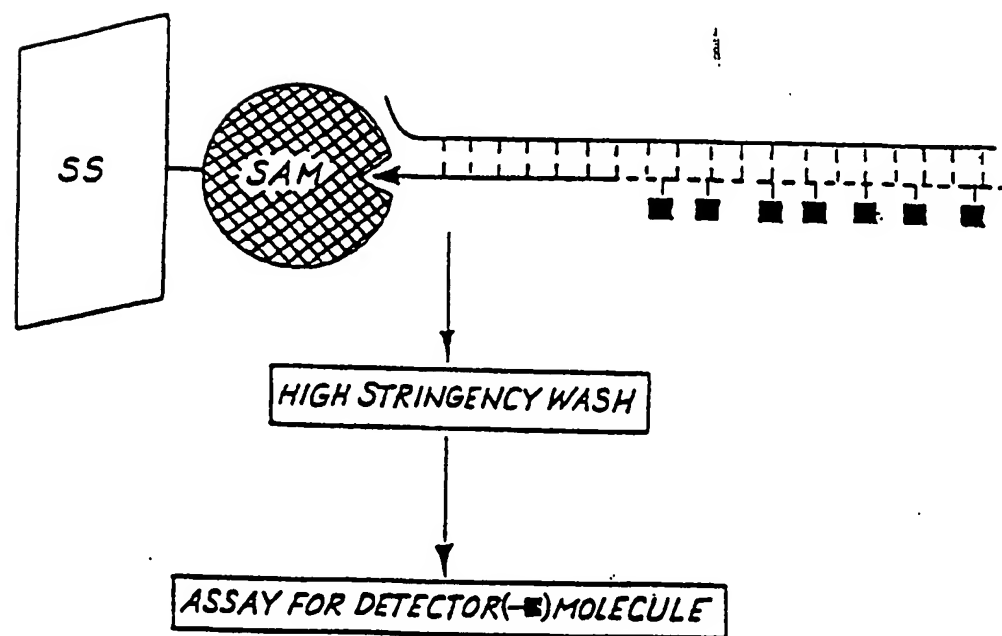
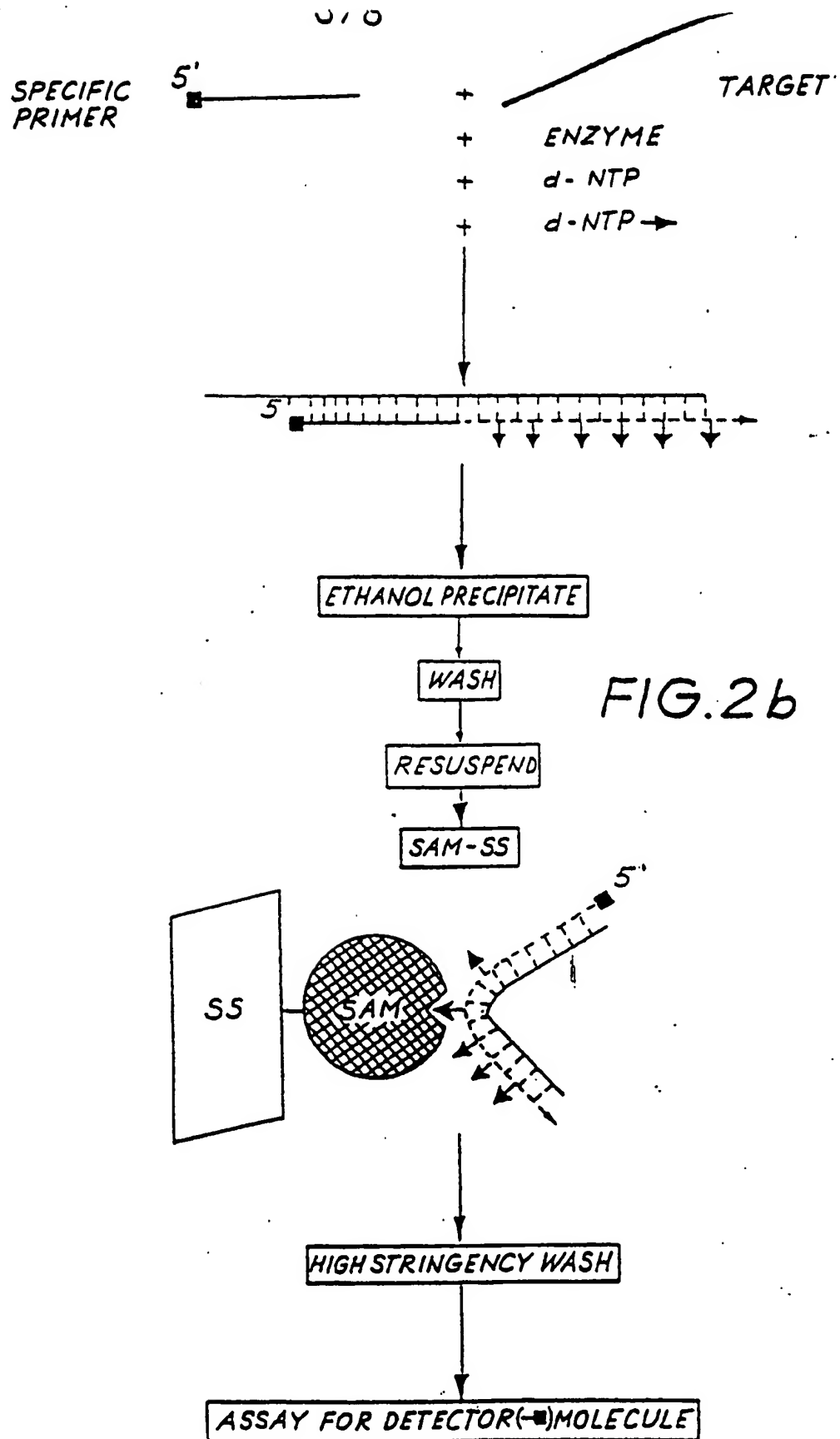
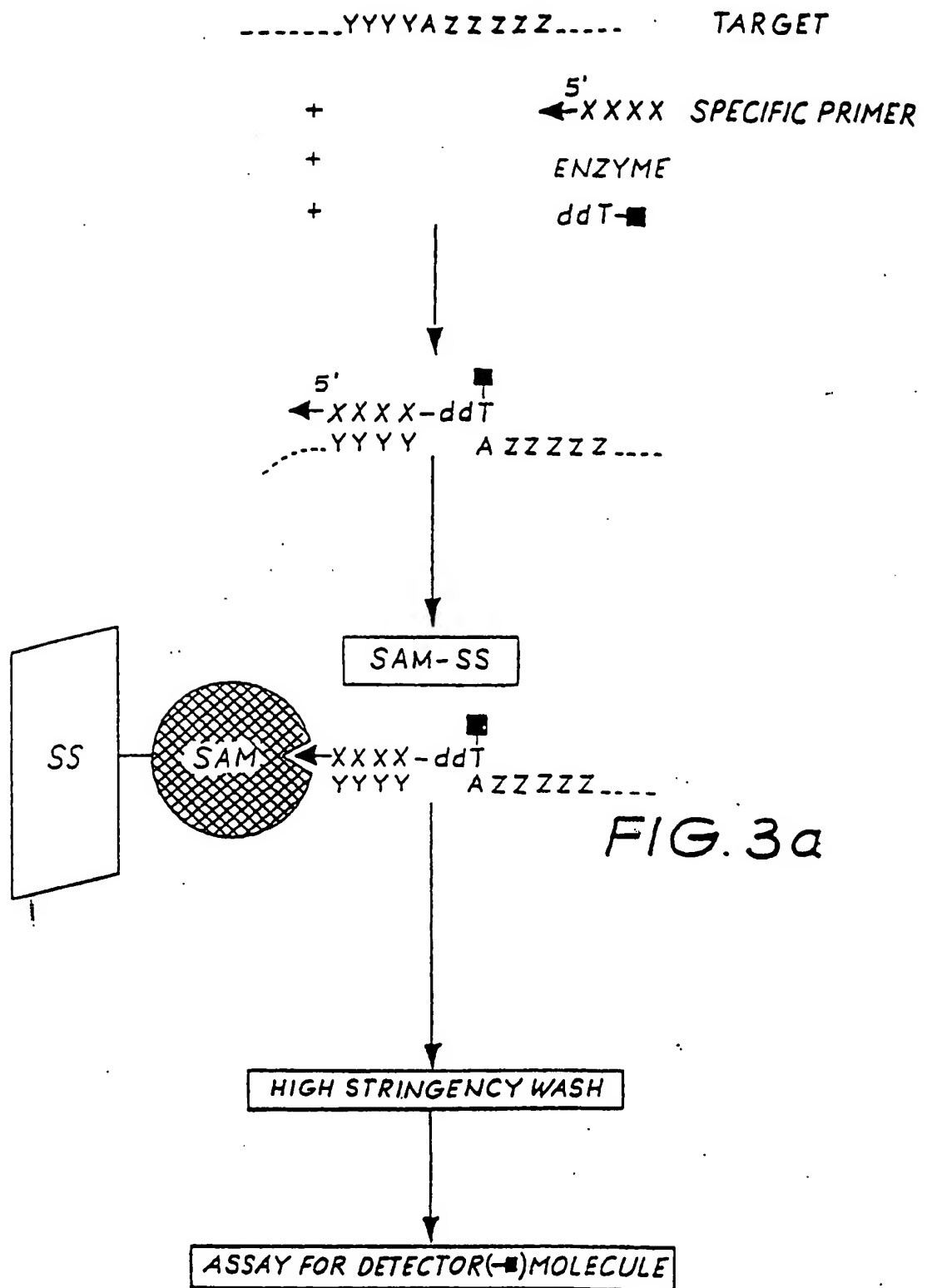
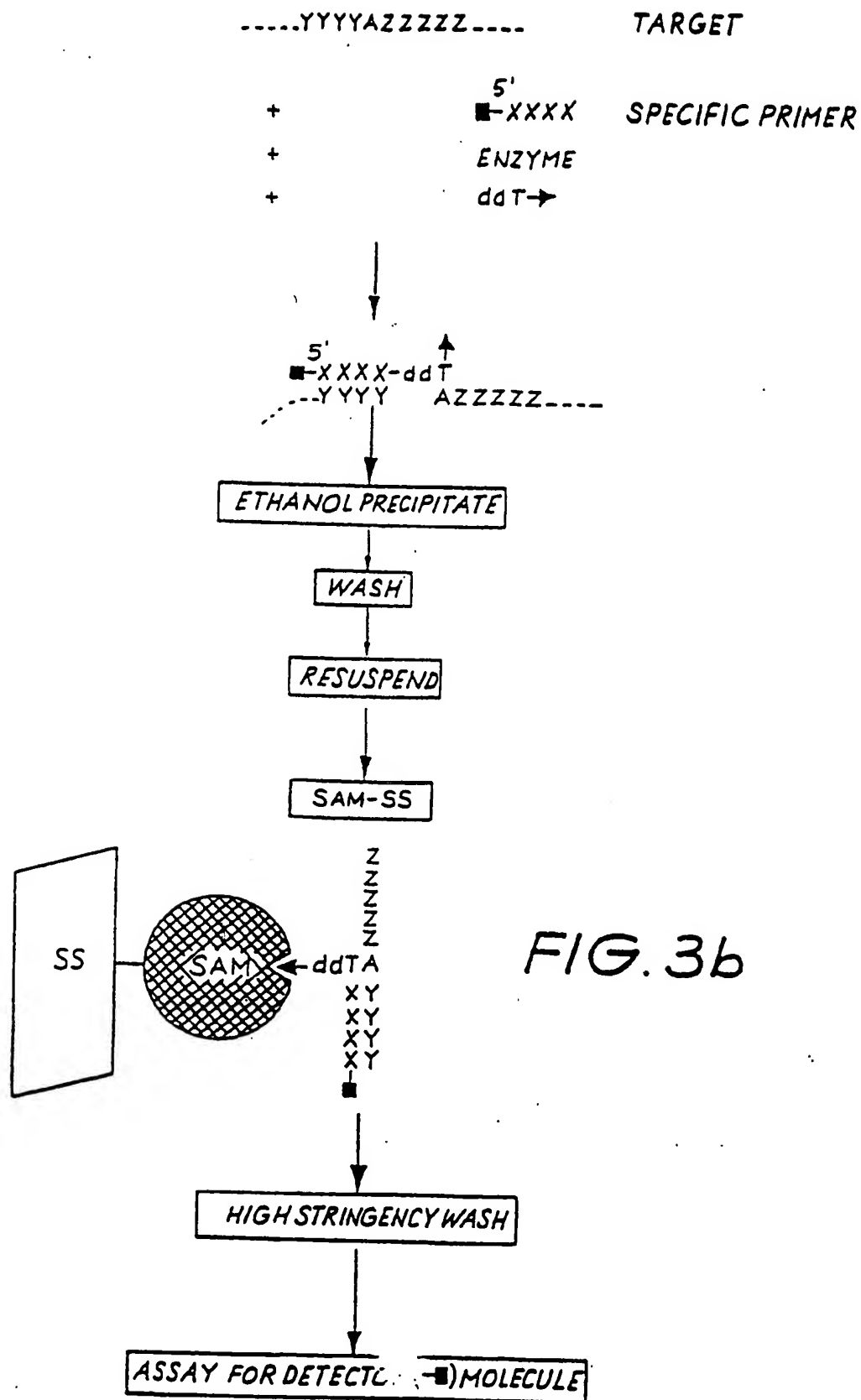


FIG.2a









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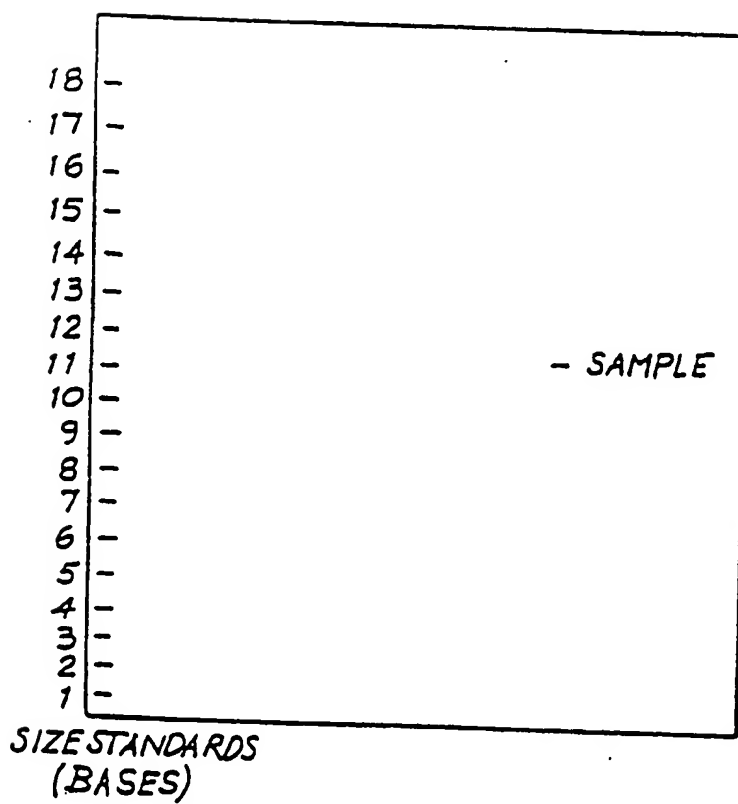
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HEAT DENATURE

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- RUN SIZE STANDARDS
- ELECTROPHORESE

FIG. 3c



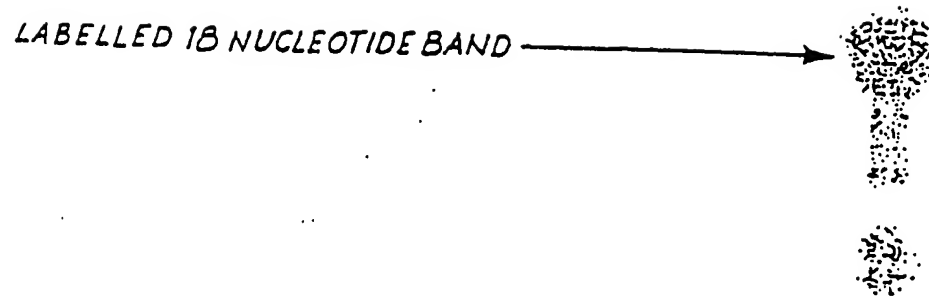


FIG. 4

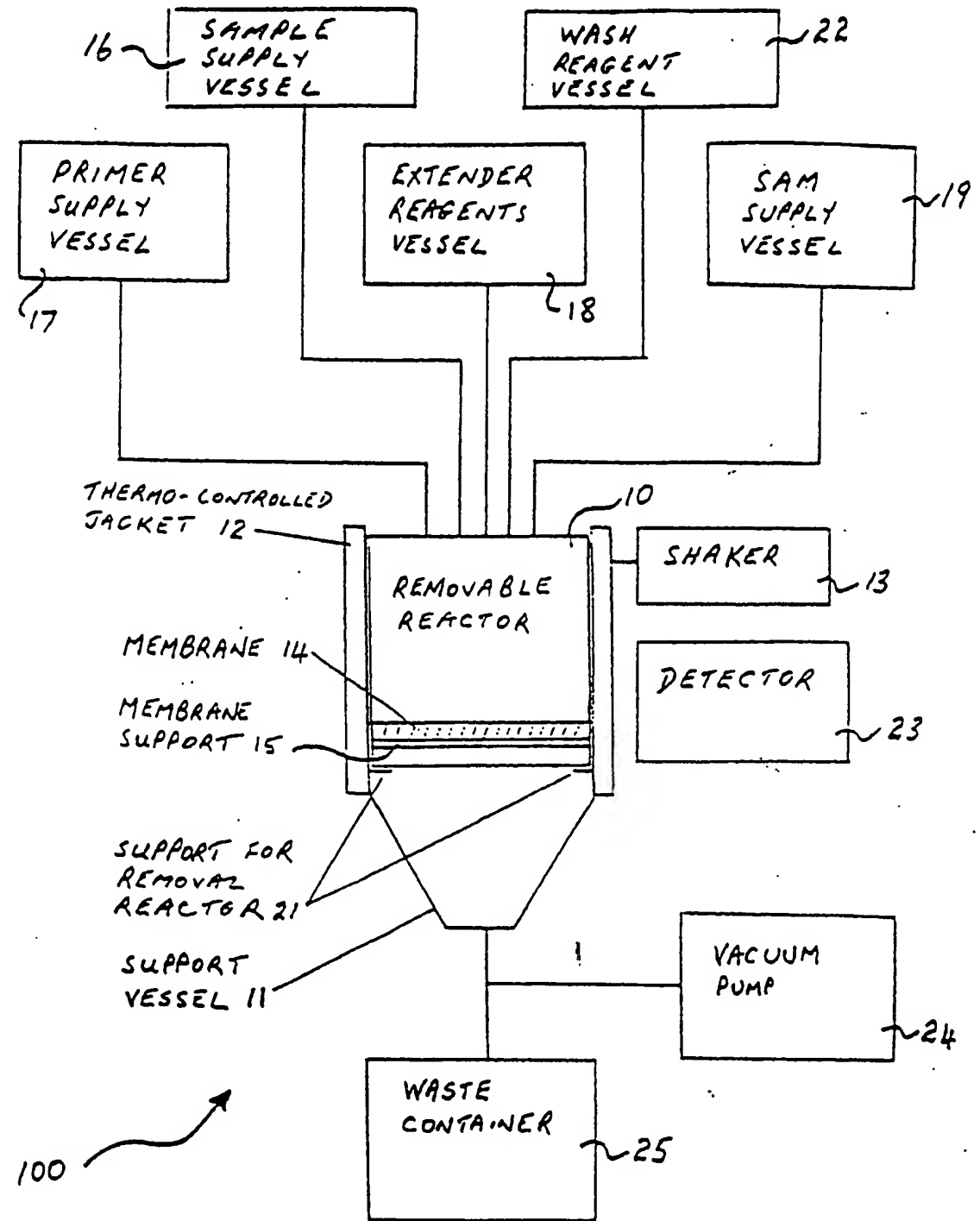


FIG. 5

I. CLASSIFICATION OF SUBJECT MATTER (if several classification symbols apply, indicate all)

According to International Patent Classification (IPC) or to both National Classification and IPC

Int. Cl. ⁵ C12Q 1/68 G01N 33/53

II. FIELDS SEARCHED

Minimum Documentation Searched 7

Classification System	Classification Symbols
IPC	C12Q 1/68

Documentation Searched other than Minimum Documentation
to the Extent that such Documents are Included in the Fields Searched 8

AU : IPC as above

Chem. Abstracts (on line): Keywords: - oligonucleotide(s), primer(s), probe(s), deoxyribonucleic acid, DNA, polymerase

III. DOCUMENTS CONSIDERED TO BE RELEVANT 9

Category	Citation of Document, with indication, where appropriate, of the relevant passages 12	Relevant to Claim No 13
P,X	US,A, 4251331 (VARY et al) 25 July 1989 (25.07.89) whole document	1,2,5,7,9,10,13, 14,17,18,21,23, 27-35,37,41
X	US,A, 4307189 (KIT) 22 December 1981 (22.12.81) whole document particularly column 2 lines 16 to 50	1,2,23-27,32,41
X	EP,A, 123513 (AMERSHAM INTERNATIONAL plc) 31 October 1984 (31.10.84) whole document	5,7,17,18,21, 23-25,32,33,41
P,X	AU,A, 30585/89 (GENENTECH INC) 27 July 1989 (27.07.89) see claims 18-20, 29, 30 and page 16 lines 16 to 35	1,9,13,23,41

(continued)

* Special categories of cited documents: 10	*T*	later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
"A" document defining the general state of the art which is not considered to be of particular relevance	"X"	document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step
"E" earlier document but published on or after the international filing date	"Y"	document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art.
"L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)	"Z"	document member of the same patent family
"O" document referring to an oral disclosure, use, exhibition or other means		
"P" document published prior to the international filing date but later than the priority date claimed		

IV. CERTIFICATION

Date of the Actual Completion of the International Search 23 May 1990 (23.05.90)	Date of Filing of this International Search Report 4 June 1990
International Searching Authority Australian Patent Office	Signature of Authorized Officer <i>S.D. Barker</i>
	S.D. BARKER

FURTHER INFORMATION CONTINUED FROM THE SECOND SHEET

<input checked="" type="checkbox"/>	AU,A, 11937/88 (ORION-YHTYMA OY) 15 September 1988 (15.09.88) whole document	1, 9, 13, 25, 27, 29-35, 41
<input checked="" type="checkbox"/>	AU,B, 62364/86 (597592) (SYNTEX INC) 12 March 1987 (12.03.87) whole document	1, 2, 9, 10, 13, 14, 26, 27, 29-35, 41

V. [] OBSERVATIONS WHERE CERTAIN CLAIMS WERE FOUND UNSEARCHABLE 1

This international search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:

1. [] Claim numbers because they relate to subject matter not required to be searched by this Authority, namely:

2. [] Claim numbers because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:

3. [] Claim numbers because they are dependent claims and are not drafted in accordance with the second and third sentences of PCT Rule 6.4 (a):

VI. [] OBSERVATIONS WHERE UNITY OF INVENTION IS LACKING 2

This International Searching Authority found multiple inventions in this international application as follows:

1. [] As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims of the international application.

2. [] As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims of the international application for which fees were paid, specifically claims:

3. [] No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claim numbers:

4. [] As all searchable claims could be searched without effort justifying an additional fee, the International Searching Authority did not invite payment of any additional fee.

Remark on Protest

[] The additional search fees were accompanied by applicant's protest.
[] No protest accompanied the payment of additional search fees.

ANNEX TO THE INTERNATIONAL SEARCH REPORT ON
INTERNATIONAL APPLICATION NO. PCT/AU 90/00058

This Annex lists the known "A" publication level patent family members relating to the patent documents cited in the above-mentioned international search report. The Australian Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

Patent Document
Cited in Search
Report

Patent Family Members

US 4851331

US 4307189

AU 30585/89 DK 4630/89 EP 359789 JP 2006724
 NO 893583 WO 8906700

AU 11937/88 DD 272664 DE 3807994 DK 1295/88
 ES 2006377 FI 880994 FR 2613077
 GB 2202328 HU 50868 IL 85480
 JP 63243875 LU 87153 NL 8800594
 NO 881064 NZ 223700 PT 86937
 SE 8800864

AU 62364/86 EP 224995 JP 62103570 US 4868104

END OF ANNEX

OTHER PUBLICATIONS

- Dean et al. (1990, Jan. 25) Nucl. Acids Res. 18(2): 345-350, "Approaches to localizing disease genes as applied to cystic fibrosis".
- Dariavach et al., "Human Ig Superfamily CTLA-4 Gene: Chromosomal Localization and Identity of Protein Sequence Between Murine and Human CTLA-4 Cytoplasmic Domains", European Journal of Immunology, vol. 18, pp. 1901-1905 (1988).
- Weber et al. Abundant Class of Human DNA Polymorphisms Which can be Typed Using the Polymerase Chain Reaction, Am. Hum. Genet., vol. 44, pp. 388-396 (1989).
- Moos et al. "Structure of Two Human Beta-Actin-related Processed Genes One of Which is Located Next to a Simple Repetitive Sequence", Embo Journal, vol. 2, No. 5, pp. 757-761, (1983).
- Chen et al., "The Human Growth Hormone Locus: Nucleotide Sequence, Biology, and Evolution", Genomic, vol 2, pp. 479-497 (1989).
- Weber et al. Dinucleotide Repeat Polymorphism at the D10S89 Locus, Nucleic Acids Research, vol. 18, No. 15, p. 4637.
- Weber et al. Dinucleotide Repeat Polymorphism at the D12S43 Locus, Nucleic Acids Research, vol. 18, No. 15, p. 4637.
- Tautz et al., Nucleic Acids Research, vol. 12, No. 10, 1984, pp. 4127-4138.
- Nakamura et al. Variable Number of Tandem Repeat (VNTR) Markers for Human Gene Mapping, Science, vol. 235, 1987, pp. 1616-1622.
- Jeffreys et al., Spontaneous Mutation Rates to New Length Alleles at Tandem-Repetitive Hypervariable Loci in Human DNA, Nature, vol. 332, 1988, pp. 278-281.
- Overhauser et al. Nucleic acids Research, vol. 15, No. 11, 1987, pp. 4617-4627.
- Jeffreys et al., Hypervariable "minisatellite" regions in Human DNA, Nature, vol. 314, 1985, pp. 67-73.
- Weber et al., Abundant Class of Human DNA Polymorphisms Which can be Typed Using the Polymerase Chain Reaction, Am. Hum. Genet., vol. 44, pp. 388-396 (1989).
- Engelke et al. Direct sequencing of enzymatically amplified human genomic DNA, Proc. Natl. Acad. U.S.A., vol. 85, pp. 544-548 (1988).
- Wong et al. Characterization of β -thalassaemia mutations using direct genomic sequencing of amplified single copy DNA, Nature, vol. 330, pp. 384-386 (1987).
- Bottstein et al., Construction of a Genetic Linkage Map in Man Using Restriction Fragment Length Polymorphisms Am. J. Hum. Genet., vol. 32, pp. 314-331 (1980).
- White et al., Chromosome Mapping with DNA Markers, Scientific American, vol. 258, pp. 40-48 (1988).
- Wallace et al. The use of synthetic oligonucleotides as hybridization probes. II Hybridization of oligonucleotides of mixed sequence to rabbit β -globin DNA, Nucl. Acid Research, vol. 9, pp. 879-984 (1981).
- Litt et al., A Hypervariable Microsatellite Revealed by In Vitro Amplification of a Dinucleotide Repeat within the Cardiac Muscle Actin Gene, Human Genet., vol. 44, pp. 397-401 (1989).